

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶ :		(1	1) International Publication Number:	WO 99/43696		
C07H 21/04, C07K 14/705, C12N 15/09, 15/63, C12Q 1/68	A1	(4	3) International Publication Date:	2 September 1999 (02.09.99)		
(21) International Application Number: PCT/US	99/038	26	(81) Designated States: AL, AM, AT	, AU, AZ, BA, BB, BG, BR,		

(22) International Filing Date: 22 February 1999 (22.02.99)

(30) Priority Data:

60/076,687 25 February 1998 (25.02.98) US 60/095,836 7 August 1998 (07.08.98) US 60/116,448 19 January 1999 (19.01.99) US

(71) Applicant: AXYS PHARMACEUTICALS, INC. [US/US]; 180 Kimball Way, South San Francisco, CA 94080 (US).

(72) Inventors: MILLER, Andrew, P.; 2131 Old Stone Mill Drive, Cranbury, NJ 08512 (US). CURRAN, Mark, Edward; 685 Poinsettia Park North, Encinitas, CA 92024 (US). HU, Ping; 3980 Via Holgura, San Diego, CA 92130 (US). RUTTER, Marc; 4559 Campus Avenue #1, San Diego, CA 92116 (US). WANG, Jian-Ying; 7478 Park Village Road, San Diego, CA 92129 (US).

(74) Agent: SHERWOOD, Pamela, J.; Bozicevic, Field & Francis LLP, Suite 200, 285 Hamilton Avenue, Palo Alto, CA 94301 (US).

PCT/US99/03826 (81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: HUMAN POTASSIUM CHANNEL GENES

(57) Abstract

Methods for isolating K+Hnov genes are provided. The K+Hnov nucleic acid compositions find use in identifying homologous or related proteins and the DNA sequences encoding such proteins; in producing compositions that modulate the expression or function of the protein; and in studying associated physiological pathways. In addition, modulation of the gene activity in vivo is used for prophylactic and therapeutic purposes, such as identification of cell type based on expression, and the like.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

ΑL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	Prance	LU	Luxembourg	SN	Senegal
ΑU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IB	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NB	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Pederation		
DE	Germany	u	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

HUMAN POTASSIUM CHANNEL GENES

INTRODUCTION

Background

5

10

15

25

30

lon channels are multi-subunit, membrane bound proteins critical for maintenance of cellular homeostasis in nearly all cell types. Channels are involved in a myriad of processes including modulation of action potentials, regulation of cardiac myocyte excitability, heart rate, vascular tone, neuronal signaling, activation and proliferation of T-cells, and insulin secretion from pancreatic islet cells. In humans, ion channels comprise extended gene families with hundreds, or perhaps thousands, of both closely related and highly divergent family members. The majority of known channels regulate the passage of sodium (Na*), chloride (Cl*), calcium (Ca**) and potassium (K*) ions across the cellular membrane.

Given their importance in maintaining normal cellular physiology, it is not surprising that ion channels have been shown to play a role in heritable human disease. Indeed, ion channel defects are involved in predisposition to epilepsy, cardiac arrhythmia (long QT syndrome), hypertension (Bartter's syndrome), cystic fibrosis, (defects in the CFTR chloride channel), several skeletal muscle disorders (hyperkalemic periodic paralysis, paramyotonia congenita, episodic ataxia) and congenital neural deafness (Jervell-Lange-Nielson syndrome), amongst others.

The potassium channel gene family is believed to be the largest and most diverse ion channel family. K* channels have critical roles in multiple cell types andpathways, and are the focus of significant investigation. Four human conditions, episodic ataxia with myokymia, long QT syndrome, epilepsy and Bartter's syndrome have been shown to be caused by defective K* ion channels. As the K* channel family is very diverse, and given that these proteins are critical components of virtually all cells, it is likely that abnormal K* channels will be involved in the etiology of additional renal, cardiovascular and central nervous system disorders of interest to the medical and pharmaceutical community.

The K* channel superfamily can be broadly classified into groups, based upon the number of transmembrane domain (TMD) segments in the mature

protein. The minK (IsK) gene contains a single TMD, and although not a channel by itself, minK associates with different K⁺ channel subunits, such as KvLQT1 and HERG to modify the activity of these channels. The inward rectifying K+ channels (GIRK, IRK, CIR, ROMK) contain 2 TMD domains and a highly conserved pore domain. Twik-1 is a member of the newly emerging 4TMD K⁺ channel subset. Twik-like channels (leak channels) are involved in maintaining the steady-state K⁺ potentials across membranes and therefore the resting potential of the cell near the equilibrium potential for potassium (Duprat *et al.* (1997) <u>EMBO J 16(17):5464-5471</u>). These proteins are particularly intriguing targets for therapeutic regulation. The 6TMD, or Shaker-like channels, presently comprise the largest subset of known K⁺ channels. The slopoke (slo) related channels, or Ca⁺⁺ regulated channels apparently have either 10 TMD, or 6 TMD with 4 additional hydrophobic domains.

10

15

20

25

30

Four transmembrane domain, tandem pore domain K+ channels (4T/2P channels) represent a new family of potassium selective ion channels involved in the control of background membrane conductances. In mammals, five channels fitting the 4T/2P architecture have been described: TWIK, TREK, TASK-1, TASK-2 and TRAAK. The 4T/2P channels all have distinct characteristics, but are all thought to be involved in maintaining the steady-state K^{\star} potentials across membranes and therefore the resting potential of the cell near the equilibrium potential for potassium (Duprat et al. (1997) EMBO J 16(17):5464-5471). These proteins are particularly intriguing targets for therapeutic regulation. Within this group, TWIK-1, TREK-1 and TASK-1 and TASK-2 are widely distributed in many different tissues, while TRAAK is present exclusively in brain, spinal cord and The 4T/2P channels have different physiologic properties; TREK-1 retina. channels, are outwardly rectifying (Fink et al. (1996) EMBO J 15(24):6854-62), while TWIK-1 channels, are inwardly rectifying (Lesage et al. (1996) EMBO J 15(5):1004-11. TASK channels are regulated by changes in PH while TRAAK channels are stimulated by arachidonic acid (Reyes et al. (1998) JBC **273**(47):30863-30869).

The degree of sequence homology between different K* channel gen s is substantial. At the amino acid level, there is about 40% similarity between

different human genes, with distinct regions having higher homology, specifically the pore domain. It has been estimated that the K+ channel gene family contains approximately 10²-10³ individual genes. Despite the large number of potential genes, an analysis of public sequence databases and the scientific literature demonstrates that only a small number, approximately 20-30, have been identified. This analysis suggests that many of these important genes remain to be identified.

Potassium channels are involved in multiple different processes and are important regulators of homeostasis in nearly all cell types. Their relevance to basic cellular physiology and role in many human diseases suggests that pharmacological agents could be designed to specific channel subtypes and these compounds then applied to a large market (Bulman, D.E. (1997) Hum Mol Genet 6:1679-1685; Ackerman, M.J. and Clapham D.E. (1997) NEJM 336:1575-1586, Curran, M.E. (1998) Current Opinion in Biotechnology 9:565-572). The variety of therapeutic agents that modulate K+ channel activity reflects the diversity of physiological roles and importance of K+ channels in cellular function. A difficulty encountered in therapeutic use of therapeutic agents that modify K+ channel activity is that the presently available compounds tend to be non-specific and elicit both positive and negative responses, thereby reducing clinical efficacy. To facilitate development of specific compounds it is desirable to have further characterize novel K+ channels for use in *in vitro* and *in vivo* assays.

Relevant Literature

20

25

30

A large body of literature exists in the general area of potassium channels. A review of the literature may be found in the series of books, "The Ion Channel Factsbook", volumes 1-4, by Edward C. Conley and William J. Brammar, Academic Press. An overview is provided of: extracellular ligand-gated ion channels (ISBN: 0121844501), intracellular ligand-gated channels (ISBN: 012184451X), Inward rectifier and intercellular channels (ISBN: 0121844528), and voltage gated channels (ISBN: 0121844536). Hille, B. (1992) "Ionic Channels of Excitabl Membranes", 2nd Ed. Sunderland MA:Sinauer Associates, also reviews potassium channels.

Jan and Jan (1997) <u>Annu. Rev. Neurosci.</u> **20**:91-123 review cloned potassium channels from eukaryotes and prokaryotes. Ackerman and Clapham (1997) <u>N. Engl. J. Med.</u> **336**:1575-1586 discuss the basic science of ion channels in connection with clinical disease. Bulman (1997) <u>Hum. Mol. Genet.</u> **6**:1679-1685 describe some phenotypic variation in ion channel disorders.

5

10

20

30

Stephan *et al.* (1994) Neurology **44**:1915-1920 describe a pedigree segregating a myotonia with muscular hypertrophy and hyperirritability as an autosomal dominant trait (rippling muscle disease, Ricker *et al.* (1989) Arch. Neurol. 46405-408). Electromyography demonstrated that mechanical stimulation provoked electrically silent contractions. The responsible gene was localized to the distal end of the long arm of chromosome 1, in a 12-cM region near D1S235.

Type II pseudohypoaldosteronism is the designation used for a syndrome of chronic mineralocorticoid-resistant hyperkalemia with hypertension. The primary abnormality in type II PHA is thought to be a specific defect of the renal secretory mechanism for potassium, which limits the kaliuretic response to, but not the sodium and chloride reabsorptive effect of, mineralocorticoid. By analysis of linkage in families with autosomal dominant transmission, Mansfield *et al.* (1997) Nature Genet. 16:202-205 demonstrated locus heterogeneity of the trait, with linkage of the PHA2 gene to 1q31-q42 and 17p11-q21.

Sequences of four transmembrane, two pore potassium channels have been previously described. Reyes et al. (1998) J Biol Chem 273(47):30863-30869 discloses a pH sensitive channel. As with the related TASK-1 and TRAAK channels, the outward rectification is lost at high external K+ concentration. The TRAAK channel is described by Fink et al. (1998) EMBO J 17(12):3297-308. A cardiac two-pore channel is described in Kim et al. (1998) Circ Res 82(4):513-8. An open rectifier potassium channel with two pore domains in tandem and having a postsynaptic density protein binding sequence at the C terminal was cloned by Leonoudakis et al. (1998) J Neurosci 18(3):868-77.

The electrophysiological properties of Task channels are of interest, (Duprat et al. (1997) EMBO J 16:5464-71). TASK currents are K+-selective, instantaneous and non-inactivating. They show an outward rectification when xternal [K+] is low, which is not observed for high [K+]out, suggesting a lack of

intrinsic voltage sensitivity. The absence of activation and inactivation kinetics as well as voltage independence are characteristic of conductances referred to as leak or background conductances. TASK is very sensitive to variations of extracellular pH in a narrow physiological range, a property probably essential for its physiological function, and suggests that small pH variations may serve a communication role in the nervous system.

SUMMARY OF THE INVENTION

10

20

25

30

Isolated nucleotide compositions and sequences are provided for *K+Hnov* genes. The *K+Hnov* nucleic acid compositions find use in identifying homologous or related genes; in producing compositions that modulate the expression or function of its encoded proteins; for gene therapy; mapping functional regions of the proteins; and in studying associated physiological pathways. In addition, modulation of the gene activity *in vivo* is used for prophylactic and therapeutic purposes, such as treatment of potassium channel defects, identification of cell type based on expression, and the like.

DESCRIPTION OF THE SPECIFIC EMBODIMENTS

Nucleic acid compositions encoding *K+Hnov* polypeptides are provided. They are used in identifying homologous or related genes; in producing compositions that modulate the expression or function of the encoded proteins; for gene therapy; mapping functional regions of the proteins; and in studying associated physiological pathways. The *K+Hnov* gene products are members of the potassium channel gene family, and have high degrees of homology to known potassium channels. The encoded polypeptides may be alpha subunits, which form the functional channel, or accessory subunits that act to modulate the channel activity.

CHARACTERIZATION OF K+HNOV

The sequence data predict that the provided K+Hnov genes encode potassium channels. Table 1 summarizes the DNA sequences, corresponding SEQ ID NOs, chromosomal locations, and polymorphisms. The provided

sequences may encode a predicted K*channel, e.g. voltage gated, inward rectifier, etc.; or a modulatory subunit.

Electrophysiologic characterization of ion channels is an important part of understanding channel function. Full length ion channel cDNAs may be combined with proper vectors to form expression constructs of each individual channel. Functional analyses of expressed channels can be performed in heterologous systems, or by expression in mammalian cell lines. For expression analyses in heterologous systems such as *Xenopus* oocytes, synthetic mRNA is made through *in vitro* transcription of each channel construct. mRNA is then injected, singly or in combination with interacting channel subunit mRNAs, into prepared oocytes and the cells allowed to express the channel for several days. Oocytes expressing the channel of interest are then analyzed by whole cell voltage clamp and patch clamp techniques.

To determine the properties of each channel when expressed in mammalian cells expression vectors specific to this type of analyses may be constructed and the resultant construct used to transform the target cells (for example human embryonic kidney (HEK) cells). Both stable and transiently expressing lines may be studied using whole cell voltage clamp and patch clamp techniques. Data obtained from EP studies includes, but is not limited to: current profiles elicited by depolarization and hyperpolarization, current-voltage (I-V) relationships, voltage dependence of activation, biophysical kinetics of channel activation, deactivation, and inactivation, reversal potential, ion selectivity, gating properties and sensitivity to channel antagonists and agonists.

15

25

Heterologous or mammalian cell lines expressing the novel channels can be used to characterize small molecules and drugs which interact with the channel. The same experiments can be used to assay for novel compounds which interact with the expressed channels.

In many cases the functional ion channel formed by K+Hnov polypeptides will be heteromultimers. Heteromultimers are known to form between different voltage gated, outward rectifying potassium channel α subunits, generally comprising four subunits, and frequently associated with auxiliary, β subunits. Typically such α subunits share a six-transmembrane domain structure (S1-S6),

with one highly positively charged domain (S4) and a pore region situated between S5 and S6. Examples of such subunits are K+Hnov4, K+Hnov9, and K+Hnov12. Channels are also formed by mutimerization of subunits of the two transmembrane and one pore architecture. It is predicted that two subunits of K+Hnov49 or K+Hnov59 will be required to form a functional channel.

5

Heteromultimers of greatest interest are those that form between subunits expressed in the same tissues, and are a combination of subunits from the same species. In addition, the formation of multimers between the subject polypeptides and subunits that form functional channels are of particular interest. The resulting channel may have decreased or increased conductance relative to a homomultimer, and may be altered in response to beta subunits or other modulatory molecules.

Known voltage gated K+ channel α subunits include Kv1.1-1.8 (Gutman *et al.* (1993) Sem. Neurosci. 5:101-106); Kv2.1-2.2; Kv3.1-3.4; Kv4.1-4.3; Kv5.1; Kv6.1; Kv7.1; Kv8.1; Kv9.1-9.2. The subunits capable of forming ion inducing channels include all of those in the Kv1 through Kv4; and Kv7 families. The Kv5.1, Kv6.1, Kv8.1 and Kv9.1-9.2 subunits may be electrically silent, but functional in modifying the properties in heteromultimers.

TABLE 1

_		_	_						_	_	_		_	_		
Channel Type	ATP-sensitive inward rectifying	Voltage gated K+ channel						Delayed rectifying K+ channel	Voltage gated K+ channel	Voltage gated K+ channel			modulatory subunit		modulatory subunit	4 transmembrane domain, 2 pore domain K+ channel
Chromosome Position	2q37	unknown						2p23	8q23	Xp21			13q14		18q12	N/A
Polymorphisms	Alternative poly(A) tail: 1236, 2395	A312C	T335C	A377G	T344C	A401G	CA410-411GG (Ala/Thr)		Alternative poly(A) tail: 2304	C321T (Pro/Leu)	A375G (Glu/Gly)	C407T (Leu/Phe)	Alternative poly(A) tail: 1427	A689G (Gly/Arg)	T365A (Ile/Asn)	N/A
Protein SEQ	SEQ ID NO:2	SEQ ID NO:4						SEQ ID NO:8	8:ON QI DES	SEQ ID NO:10			SEQ ID NO:12		SEQ ID NO:14	SEQ ID NO:16
cDNA SEQ	SEQ ID NO:1	SEQ ID NO:3						SEQ ID NO:5	SEQ ID NO:7	SEQ ID NO:9			SEQ ID NO:11		SEQ ID NO:13	SEQ ID NO:15
Name	K+Hnov1	K+Hnov4						K+Hnov6	K+Hnov9	K+Hnov12			K+Hnov15		K+Hnov27	K+Hnov2

K+Hnov 11	SEQ ID NO:17	SEQ ID NO:18	N/A	K/A	Human ortholog of murine gene, 6
					transmembrane dominas, voltage gated, delayed rectifier K+ channel
K+Hnov 14	SEQ ID NO:19	SEQ ID NO:20	C3168T	12q14	6 transmembrane domain, voltage gated K+ channel
K+Hnov28	SEQ ID NO:21-24	SEQ ID NO:25	4 alternative 5' splices	3q29	Modulatory subunit
K+Hnov42	SEQ ID NO:26	SEQ ID NO:27	G1162A; T1460A; T2496A	8q11	Homology to K+ channel protein of C. elegans
K+Hnov44	SEQ ID NO:28-29	SEQ ID NO:30	N/A	22p13	beta-subunit.
K'Hnov49	SEQ ID NO:80	SEQ ID NO:81	(ATCT), repeats in the 3' UTR sequence, starting at position 2186	1941	4T/2P channel; linked to the disease loci for rippling muscle disease 1 (RMD1), and type II pseudohypoaldosteronism
K*Hnov59	SEQ ID NO:82	SEQ ID NO:83	N/A	chr19	4T/2P channel

K+HNOV NUCLEIC ACID COMPOSITIONS

As used herein, the term "K+Hnov" is generically used to refer to any one of the provided genetic sequences listed in Table 1. Where a specific K+Hnov sequence is intended, the numerical designation, e.g. K49 or K59, will be added. Nucleic acids encoding K+Hnov potassium channels may be cDNA or genomic DNA or a fragment thereof. The term "K+Hnov gene" shall be intended to mean the open reading frame encoding any of the provided K+Hnov polypeptides, introns, as well as adjacent 5' and 3' non-coding nucleotide sequences involved in the regulation of expression, up to about 20 kb beyond the coding region, but possibly further in either direction. The gene may be introduced into an appropriate vector for extrachromosomal maintenance or for integration into a host genome.

The term "cDNA" as used herein is intended to include all nucleic acids that share the arrangement of sequence elements found in native mature mRNA species, where sequence elements are exons and 3' and 5' non-coding regions. Normally mRNA species have contiguous exons, with the intervening introns, when present, removed by nuclear RNA splicing, to create a continuous open reading frame encoding a K+Hnov protein.

A genomic sequence of interest comprises the nucleic acid present between the initiation codon and the stop codon, as defined in the listed sequences, including all of the introns that are normally present in a native chromosome. It may further include the 3' and 5' untranslated regions found in the mature mRNA. It may further include specific transcriptional and translational regulatory sequences, such as promoters, enhancers, etc., including about 1 kb, but possibly more, of flanking genomic DNA at either the 5' or 3' end of the transcribed region. The genomic DNA may be isolated as a fragment of 100 kbp or smaller; and substantially free of flanking chromosomal sequence. The genomic DNA flanking the coding region, either 3' or 5', or internal regulatory sequences as sometimes found in introns, contains sequences required for proper tissue and stage specific expression.

20

25

30

The sequence of the 5' flanking region may be utilized for promoter elements, including enhancer binding sites, that provide for developmental regulation in tissues where *K+Hnov* genes are expressed. The tissue specific expression is useful for determining the pattern of expression, and for providing promoters that mimic the native pattern of expression. Naturally occurring polymorphisms in the promoter regions are useful for determining natural variations in expression, particularly those that may be associated with disease.

Alternatively, mutations may be introduced into the promoter regions to determine the effect of altering expression in experimentally defined systems. Methods for the identification of specific DNA motifs involved in the binding of transcriptional factors are known in the art, e.g. sequence similarity to known binding motifs, gel retardation studies, etc. For examples, see Blackwell et al. (1995) Mol Med 1: 194-205; Mortlock et al. (1996) Genome Res. 6: 327-33; and Joulin and Richard-Foy (1995) Eur J Biochem 232: 620-626.

The regulatory sequences may be used to identify *cis* acting sequences required for transcriptional or translational regulation of *K+Hnov* expression, especially in different tissues or stages of development, and to identify *cis* acting sequences and *trans* acting factors that regulate or mediate *K+Hnov* expression. Such transcription or translational control regions may be operably linked to a *K+Hnov* gene in order to promote expression of wild type or altered *K+Hnov* or other proteins of interest in cultured cells, or in embryonic, fetal or adult tissues, and for gene therapy.

15

20

The nucleic acid compositions of the subject invention may encode all or a part of the subject polypeptides. Double or single stranded fragments may be obtained of the DNA sequence by chemically synthesizing oligonucleotides in accordance with conventional methods, by restriction enzyme digestion, by PCR amplification, etc. For the most part, DNA fragments will be of at least 15 nt, usually at least 18 nt or 25 nt, and may be at least about 50 nt. Such small DNA fragments are useful as primers for PCR, hybridization screening probes, etc. Larger DNA fragments, i.e. greater than 100 nt are useful for production of the ncoded polypeptide. For use in amplification reactions, such as PCR, a pair of

primers will b used. The exact composition of the primer sequences is not critical to the invention, but for most applications the primers will hybridize to the subject sequence under stringent conditions, as known in the art. It is preferable to choose a pair of primers that will generate an amplification product of at least about 50 nt, preferably at least about 100 nt. Algorithms for the selection of primer sequences are generally known, and are available in commercial software packages. Amplification primers hybridize to complementary strands of DNA, and will prime towards each other.

The K+Hnov genes are isolated and obtained in substantial purity, generally as other than an intact chromosome. Usually, the DNA will be obtained substantially free of other nucleic acid sequences that do not include a K+Hnov sequence or fragment thereof, generally being at least about 50%, usually at least about 90% pure and are typically "recombinant", i.e. flanked by one or more nucleotides with which it is not normally associated on a naturally occurring chromosome.

15

The DNA may also be used to identify expression of the gene in a biological specimen. The manner in which one probes cells for the presence of particular nucleotide sequences, as genomic DNA or RNA, is well established in the literature and does not require elaboration here. DNA or mRNA is isolated from a cell sample. The mRNA may be amplified by RT-PCR, using reverse transcriptase to form a complementary DNA strand, followed by polymerase chain reaction amplification using primers specific for the subject DNA sequences. Alternatively, the mRNA sample is separated by gel electrophoresis, transferred to a suitable support, e.g. nitrocellulose, nylon, etc., and then probed with a fragment of the subject DNA as a probe. Other techniques, such as oligonucleotide ligation assays, in situ hybridizations, and hybridization to DNA probes arrayed on a solid chip may also find use. Detection of mRNA hybridizing to the subject sequence is indicative of K+Hnov gene expression in the sample.

The sequence of a K+Hnov gene, including flanking promoter regions and coding regions, may be mutated in various ways known in the art to gen rate targ ted changes in promoter strength, sequence of the encoded prot in, etc.

The DNA sequence or protein product of such a mutation will usually be substantially similar to the sequences provided herein, *i.e.* will differ by at least one nucleotide or amino acid, respectively, and may differ by at least two but not more than about ten nucleotides or amino acids. The sequence changes may be substitutions, insertions or deletions. Deletions may further include larger changes, such as deletions of a domain or exon. Other modifications of interest include epitope tagging, e.g. with the FLAG system, HA, etc. For studies of subcellular localization, fusion proteins with green fluorescent proteins (GFP) may be used.

10

20

Techniques for *in vitro* mutagenesis of cloned genes are known. Examples of protocols for site specific mutagenesis may be found in Gustin *et al.*, *Biotechniques* 14:22 (1993); Barany, *Gene* 37:111-23 (1985); Colicelli *et al.*, *Mol Gen Genet* 199:537-9 (1985); and Prentki *et al.*, *Gene* 29:303-13 (1984). Methods for site specific mutagenesis can be found in Sambrook *et al.*, *Molecular Cloning: A Laboratory Manual*, CSH Press 1989, pp. 15.3-15.108; Weiner *et al.*, *Gene* 126:35-41 (1993); Sayers *et al.*, *Biotechniques* 13:592-6 (1992); Jones and Winistorfer, *Biotechniques* 12:528-30 (1992); Barton *et al.*, *Nucleic Acids Res* 18:7349-55 (1990); Marotti and Tomich, *Gene Anal Tech* 6:67-70 (1989); and Zhu, *Anal Biochem* 177:120-4 (1989). Such mutated genes may be used to study structure-function relationships of *K+Hnov*, or to alter properties of the protein that affect its function or regulation.

Homologs and orthologs of K+Hnov genes are identified by any of a number of methods. A fragment of the provided cDNA may be used as a hybridization probe against a cDNA library from the target organism of interest, where low stringency conditions are used. The probe may be a large fragment, or one or more short degenerate primers. Nucleic acids having sequence similarity are detected by hybridization under low stringency conditions, for example, at 50°C and 6XSSC (0.9 M sodium chloride/0.09 M sodium citrate) and remain bound when subjected to washing at 55°C in 1XSSC (0.15 M sodium chloride/0.015 M sodium citrate). Sequence identity may be determined by hybridization under stringent conditions, for example, at 50°C or higher and

0.1XSSC (15 mM sodium chloride/01.5 mM sodium citrate). Nucleic acids having a region of substantial identity to the provided K+Hnov sequences, e.g. allelic variants, genetically altered versions of the gene, etc., bind to the provided K+Hnov sequences under stringent hybridization conditions. By using probes, particularly labeled probes of DNA sequences, one can isolate homologous or related genes. The source of homologous genes may be any species, e.g. primate species, particularly human; rodents, such as rats and mice, canines, felines, bovines, ovines, equines, yeast, nematodes, etc.

10

20

25

Between mammalian species, e.g. human and mouse, homologs have substantial sequence similarity, i.e. at least 75% sequence identity between nucleotide sequences, in some cases 80 or 90% sequence identity, and may be as high as 95% sequence identity between closely related species. Sequence similarity is calculated based on a reference sequence, which may be a subset of a larger sequence, such as a conserved motif, coding region, flanking region, etc. A reference sequence will usually be at least about 18 nt long, more usually at least about 30 nt long, and may extend to the complete sequence that is being compared. Algorithms for sequence analysis are known in the art, such as BLAST, described in Altschul et al. (1990), J. Mol. Biol. 215:403-10. In general, variants of the invention have a sequence identity greater than at least about 65%, preferably at least about 75%, more preferably at least about 85%, and may be greater than at least about 90% or more as determined by the Smith-Waterman homology search algorithm as implemented in MPSRCH program (Oxford Molecular). Exemplary search parameters for use with the MPSRCH program in order to identify sequences of a desired sequence identity are as follows: gap open penalty: 12; and gap extension penalty: 1.

K+HNOV POLYPEPTIDES

The subject nucleic acid sequences may be employed for producing all or portions of K+Hnov polypeptides. For expression, an expression cassette may be employed. The expression vector will provide a transcriptional and translational initiation region, which may be inducible or constitutive, where the coding region

is operably linked under the transcriptional control of the transcriptional initiation region, and a transcriptional and translational termination region. These control regions may be native to a *K+Hnov* gene, or may be derived from exogenous sources.

5

10

The peptide may be expressed in prokaryotes or eukaryotes in accordance with conventional ways, depending upon the purpose for expression. For large scale production of the protein, a unicellular organism, such as *E. coli, B. subtilis, S. cerevisiae*, insect cells in combination with baculovirus vectors, or cells of a higher organism such as vertebrates, particularly mammals, *e.g.* COS 7 cells, may be used as the expression host cells. In some situations, it is desirable to express the *K+Hnov* gene in eukaryotic cells, where the *K+Hnov* protein will benefit from native folding and post-translational modifications. Small peptides can also be synthesized in the laboratory. Peptides that are subsets of the complete *K+Hnov* sequence may be used to identify and investigate parts of the protein important for function, or to raise antibodies directed against these regions.

Fragments of interest include the transmembrane and pore domains, the signal sequences, regions of interaction between subunits, *etc.* Such domains will usually include at least about 20 amino acids of the provided sequence, more usually at least about 50 amino acids, and may include 100 amino acids or more, up to the complete domain. Binding contacts may be comprised of noncontiguous sequences, which are brought into proximity by the tertiary structure of the protein. The sequence of such fragments may be modified through manipulation of the coding sequence, as described above. Truncations may be performed at the carboxy or amino terminus of the fragment, e.g. to determine the minimum sequence required for biological activity.

With the availability of the protein or fragments thereof in large amounts, by employing an expression host, the protein may be isolated and purified in accordance with conventional ways. A lysate may be prepared of the expression host and the lysate purified using HPLC, exclusion chromatography, gel electrophoresis, affinity chromatography, or oth r purification technique. The

purified protein will generally be at least about 80% pure, preferably at least about 90% pure, and may be up to and including 100% pure. Pure is intended to mean free of other proteins, as well as cellular debris.

The expressed K+Hnov polypeptides are useful for the production of antibodies, where short fragments provide for antibodies specific for the particular polypeptide, and larger fragments or the entire protein allow for the production of antibodies over the surface of the polypeptide. Antibodies may be raised to the wild-type or variant forms of K+Hnov. Antibodies may be raised to isolated peptides corresponding to specific domains, e.g. the pore domain and the transmembrane domain, or to the native protein.

10

20

25

Antibodies are prepared in accordance with conventional ways, where the expressed polypeptide or protein is used as an immunogen, by itself or conjugated to known immunogenic carriers, e.g. KLH, pre-S HBsAg, other viral or eukaryotic proteins, or the like. Various adjuvants may be employed, with a series of injections, as appropriate. For monoclonal antibodies, after one or more booster injections, the spleen is isolated, the lymphocytes immortalized by cell fusion, and then screened for high affinity antibody binding. The immortalized cells, i.e. hybridomas, producing the desired antibodies may then be expanded. For further description, see Monoclonal Antibodies: A Laboratory Manual, Harlow and Lane eds., Cold Spring Harbor Laboratories, Cold Spring Harbor, New York, 1988. If desired, the mRNA encoding the heavy and light chains may be isolated and mutagenized by cloning in E. coli, and the heavy and light chains mixed to further enhance the affinity of the antibody. Alternatives to in vivo immunization as a method of raising antibodies include binding to phage "display" libraries, usually in conjunction with in vitro affinity maturation.

K+HNOV GENOTYPING

The subject nucleic acid and/or polypeptide compositions may be used to genotyping and other analysis for the presence of polymorphisms in the sequence, or variation in the expression of the subject genes. Genotyping may b performed to det rmine whether a particular polymorphisms is associated with

a disease state or genetic predisposition to a disease state, particularly diseases associated with defects in excitatory properties of cells, e.g. cardiac, muscle, renal and neural cells. Disease of interest include rippling muscle disease, and type II psuedohypoaldosteronism.

5

15

20

25

30

Clinical disorders associated with K+ channel defects include long-QT syndrome; a congenital disorder affecting 1 in 10,000-15,000. Affected individuals have a prolonged QT interval in the electrocardiogram due to a delayed repolarization of the ventricle. Genetic linkage analyses identified two loci for long QT syndrome, LQT1, in 11p15.5 and LQT2, in 7q35-36. Positional cloning techniques identified the novel K+ channel KvLQT1 on chromosome 11 while candidate gene analysis identified causative mutations in the HERG K+ channel for LQT2.

The weaver mouse exhibits several abnormal neurological symptoms, including severe ataxia, loss of granule cell neurons in the cerebellum and dopaminergic cells in the substantia nigra, as well as seizures and male infertility. A G-protein-coupled K+ channel having a mutation in the conserved pore domain has been determined to cause the disease. The pancreatic-islet \(\mathcal{G}\)-cell ATP-sensitive K+ channel (KATP) is composed of two subunits, the sulfonylurea receptor (SUR) and the inward rectifier K+ channel Kir6.2. Mutations in both SUR and Kir6.2 have been identified in patients with persistent hyperinsulinemic hypoglycemia of infancy, which is caused by unregulated secretion of insulin.

Genotyping may also be performed for pharmacogenetic analysis to assess the association between an individual's genotype and that individual's ability to react to a therapeutic agent. Differences in target sensitivity can lead to toxicity or therapeutic failure. Relationships between polymorphisms in channel expression or specificity can be used to optimize therapeutic dose administration.

Genetic polymorphisms are identified in the K+Hnov gene (examples are listed in table 1), e.g. the repeat variation in the 3' UTR of K49. Nucleic acids comprising the polymorphic sequences are used to screen patients for altered reactivity and adverse side effects in response to drugs that act on K+ channels.

K+Hnov genotyping is performed by DNA or RNA sequence and/or hybridization analysis of any convenient sample from a patient, e.g. biopsy material, blood sample, scrapings from cheek, etc. A nucleic acid sample from an individual is analyzed for the presence of polymorphisms in K+Hnov, particularly those that affect the activity, responsiveness or expression of K+Hnov. Specific sequences of interest include any polymorphism that leads to changes in basal expression in one or more tissues, to changes in the modulation of K+Hnov expression, or alterations in K+Hnov specificity and/or activity.

The effect of a polymorphism in K+Hnov gene sequence on the response to a particular agent may be determined by *in vitro* or *in vivo* assays. Such assays may include monitoring during clinical trials, testing on genetically defined cell lines, etc. The response of an individual to the agent can then be predicted by determining the K+Hnov genotype with respect to the polymorphism. Where there is a differential distribution of a polymorphism by racial background, guidelines for drug administration can be generally tailored to a particular ethnic group.

10

20

25

Biochemical studies may be performed to determine whether a sequence polymorphism in a *K+Hnov* coding region or control regions is associated with disease, for example the association of K+Hnov 9 with idiopathic generalized epilepsy. Disease associated polymorphisms may include deletion or truncation of the gene, mutations that alter expression level, that affect the electrical activity of the channel, *etc.*

A number of methods are available for analyzing nucleic acids for the presence of a specific sequence. Where large amounts of DNA are available, genomic DNA is used directly. Alternatively, the region of interest is cloned into a suitable vector and grown in sufficient quantity for analysis. The nucleic acid may be amplified by conventional techniques, such as the polymerase chain reaction (PCR), to provide sufficient amounts for analysis. The use of the polymerase chain reaction is described in Saiki et al. (1985) Science 239:487, and a review of current techniques may be found in Sambrook et al. Molecular Cloning: A Laboratory Manual, CSH Press 1989, pp.14.2–14.33. Amplification may b used

to determine whether a polymorphism is present, by using a primer that is specific for the polymorphism. Alternatively, various methods are known in the art that utilize oligonucleotide ligation as a means of detecting polymorphisms, for examples see Riley *et al.* (1990) N.A.R. 18:2887-2890; and Delahunty *et al.* (1996) Am. J. Hum. Genet. 58:1239-1246.

5

10

15

20

25

A detectable label may be included in an amplification reaction. Suitable labels include fluorochromes, e.g. fluorescein isothiocyanate (FITC), rhodamine, Texas Red, phycoerythrin, allophycocyanin, 6-carboxyfluorescein (6-FAM), 2',7'-dimethoxy-4',5'- dichloro-6-carboxyfluorescein (JOE), 6-carboxy-X-rhodamine (ROX), 6-carboxy-2',4',7',4,7- hexachlorofluorescein (HEX), 5-carboxyfluorescein (5-FAM) or N,N,N',N'-tetramethyl-6- carboxyrhodamine (TAMRA), radioactive labels, e.g. 32P, 35S, 3H; etc. The label may be a two stage system, where the amplified DNA is conjugated to biotin, haptens, etc. having a high affinity binding partner, e.g. avidin, specific antibodies, etc., where the binding partner is conjugated to a detectable label. The label may be conjugated to one or both of the primers. Alternatively, the pool of nucleotides used in the amplification is labeled, so as to incorporate the label into the amplification product.

The sample nucleic acid, e.g. amplified or cloned fragment, is analyzed by one of a number of methods known in the art. The nucleic acid may be sequenced by dideoxy or other methods. Hybridization with the variant sequence may also be used to determine its presence, by Southern blots, dot blots, etc. The hybridization pattern of a control and variant sequence to an array of oligonucleotide probes immobilised on a solid support, as described in U.S. 5,445,934, or in WO95/35505, may also be used as a means of detecting the presence of variant sequences. Single strand conformational polymorphism (SSCP) analysis, denaturing gradient gel electrophoresis (DGGE), mismatch cleavage detection, and heteroduplex analysis in gel matrices are used to detect conformational changes created by DNA sequence variation as alterations in electrophoretic mobility. Alternatively, where a polymorphism creates or destroys a recognition site for a restriction endonuclease (restriction fragment length polymorphism, RFLP), the sample is digested with that endonuclease, and the

products size fractionated to determine whether the fragment was digested. Fractionation is performed by gel or capillary electrophoresis, particularly acrylamide or agarose gels.

In one embodiment of the invention, an array of oligonucleotides are provided, where discrete positions on the array are complementary to one or more of the provided sequences, e.g. oligonucleotides of at least 12 nt, frequently 20 nt, or larger, and including the sequence flanking a polymorphic position in a K*Hnov sequence; coding sequences for different K*Hnov channels, panels of ion channels comprising one or more of the provided K* channels; etc. Such an array may comprise a series of oligonucleotides, each of which can specifically hybridize to a different polymorphism. For examples of arrays, see Hacia et al. (1996) Nature Genetics 14:441-447; Lockhart et al. (1996) Nature Biotechnol. 14:1675-1680; and De Risi et al. (1996) Nature Genetics 14:457-460.

10

15

20

25

30

Screening for polymorphisms in K+Hnov may be based on the functional or antigenic characteristics of the protein. Protein truncation assays are useful in detecting deletions that may affect the biological activity of the protein. Various immunoassays designed to detect polymorphisms in K+Hnov proteins may be used in screening. Where many diverse genetic mutations lead to a particular disease phenotype, functional protein assays have proven to be effective screening tools. The activity of the encoded K+Hnov protein as a potassium channel may be determined by comparison with the wild-type protein.

Antibodies specific for a K+Hnov may be used in staining or in immunoassays. Samples, as used herein, include biological fluids such as semen, blood, cerebrospinal fluid, tears, saliva, lymph, dialysis fluid and the like; organ or tissue culture derived fluids; and fluids extracted from physiological tissues. Also included in the term are derivatives and fractions of such fluids. The cells may be dissociated, in the case of solid tissues, or tissue sections may be analyzed. Alternatively a lysate of the cells may be prepared.

Diagnosis may be performed by a number of methods to determine the absence or presence or altered amounts of normal or abnormal K+Hnov polypeptides in patient cells. For exampl, detection may utilize staining of cells

or histological sections, performed in accordance with conventional methods. The antibodies of interest are added to the cell sample, and incubated for a period of time sufficient to allow binding to the epitope, usually at least about 10 minutes. The antibody may be labeled with radioisotopes, enzymes, fluorescers, chemiluminescers, or other labels for direct detection. Alternatively, a second stage antibody or reagent is used to amplify the signal. Such reagents are well known in the art. For example, the primary antibody may be conjugated to biotin, with horseradish peroxidase-conjugated avidin added as a second stage reagent. Alternatively, the secondary antibody conjugated to a flourescent compound, e.g. flourescein, rhodamine, Texas red, etc. Final detection uses a substrate that undergoes a color change in the presence of the peroxidase. The absence or presence of antibody binding may be determined by various methods, including flow cytometry of dissociated cells, microscopy, radiography, scintillation counting, etc.

15

20

30

MODULATION OF GENE EXPRESSION

The K+Hnov genes, gene fragments, or the encoded protein or protein fragments are useful in gene therapy to treat disorders associated with K+Hnov defects. Expression vectors may be used to introduce the K+Hnov gene into a cell. Such vectors generally have convenient restriction sites located near the promoter sequence to provide for the insertion of nucleic acid sequences. Transcription cassettes may be prepared comprising a transcription initiation region, the target gene or fragment thereof, and a transcriptional termination region. The transcription cassettes may be introduced into a variety of vectors, e.g. plasmid; retrovirus, e.g. lentivirus; adenovirus; and the like, where the vectors are able to transiently or stably be maintained in the cells, usually for a period of at least about one day, more usually for a period of at least about several days to several weeks.

The gene or K+Hnov protein may be introduced into tissues or host cells by any number of routes, including viral infection, microinjection, or fusion of v sicles. Jet injection may also b used for intramuscular administration, as

described by Furth et al. (1992) Anal Biochem 205:365-368. The DNA may be coated onto gold microparticles, and delivered intradermally by a particle bombardment device, or "gene gun" as described in the literature (see, for example, Tang et al. (1992) Nature 356:152-154), where gold microprojectiles are coated with the K+Hnov or DNA, then bombarded into skin cells.

5

25

30

Antisense molecules can be used to down-regulate expression of K+Hnov in cells. The anti-sense reagent may be antisense oligonucleotides (ODN), particularly synthetic ODN having chemical modifications from native nucleic acids, or nucleic acid constructs that express such anti-sense molecules as RNA. The antisense sequence is complementary to the mRNA of the targeted gene, and inhibits expression of the targeted gene products. Antisense molecules inhibit gene expression through various mechanisms, e.g. by reducing the amount of mRNA available for translation, through activation of RNAse H, or steric hindrance. One or a combination of antisense molecules may be administered, where a combination may comprise multiple different sequences.

Antisense molecules may be produced by expression of all or a part of the target gene sequence in an appropriate vector, where the transcriptional initiation is oriented such that an antisense strand is produced as an RNA molecule. Alternatively, the antisense molecule is a synthetic oligonucleotide. Antisense oligonucleotides will generally be at least about 7, usually at least about 12, more usually at least about 20 nucleotides in length, and not more than about 500, usually not more than about 50, more usually not more than about 35 nucleotides in length, where the length is governed by efficiency of inhibition, specificity, including absence of cross-reactivity, and the like. It has been found that short oligonucleotides, of from 7 to 8 bases in length, can be strong and selective inhibitors of gene expression (see Wagner *et al.* (1996) Nature Biotechnology 14:840-844).

A specific region or regions of the endogenous sense strand mRNA sequence is chosen to be complemented by the antisense sequence. Selection of a specific sequence for the oligonucleotide may us an empirical method, where several candidate sequences are assayed for inhibition of expression of

the target gene in an *in vitro* or animal model. A combination of sequences may also be used, where several regions of the mRNA sequence are selected for antisense complementation.

Antisense oligonucleotides may be chemically synthesized by methods known in the art (see Wagner et al. (1993) supra. and Milligan et al., supra.) Preferred oligonucleotides are chemically modified from the native phosphodiester structure, in order to increase their intracellular stability and binding affinity. A number of such modifications have been described in the literature, which alter the chemistry of the backbone, sugars or heterocyclic bases.

5

10

15

20

25

30

Among useful changes in the backbone chemistry are phosphorothioates; phosphorodithioates, where both of the non-bridging oxygens are substituted with sulfur; phosphoroamidites; alkyl phosphotriesters and boranophosphates. Achiral 3'-S-5'-O-3'-O'-5'-S-phosphorothioate, derivatives include phosphate phosphorothioate, 3'-CH2-5'-O-phosphonate and 3'-NH-5'-O-phosphoroamidate. Peptide nucleic acids replace the entire ribose phosphodiester backbone with a peptide linkage. Sugar modifications are also used to enhance stability and affinity. The α-anomer of deoxyribose may be used, where the base is inverted with respect to the natural β-anomer. The 2'-OH of the ribose sugar may be altered to form 2'-O-methyl or 2'-O-allyl sugars, which provides resistance to degradation without comprising affinity. Modification of the heterocyclic bases Some useful substitutions include must maintain proper base pairing. deoxyuridine for deoxythymidine; 5-methyl-2'-deoxycytidine and 5-bromo-2'deoxycytidine for deoxycytidine. 5- propynyl-2'-deoxyuridine and 5-propynyl-2'deoxycytidine have been shown to increase affinity and biological activity when substituted for deoxythymidine and deoxycytidine, respectively.

As an alternative to anti-sense inhibitors, catalytic nucleic acid compounds, e.g. ribozymes, anti-sense conjugates, etc. may be used to inhibit gene expression. Ribozymes may be synthesized in vitro and administered to the patient, or may be needed on an expression v ctor, from which the ribozyme is synthesized in the targeted cell (for example, see International patent application)

WO 9523225, and Beigelman et al. (1995) <u>Nucl. Acids Res</u> 23:4434-42). Examples of oligonucleotides with catalytic activity are described in WO 9506764. Conjugates of anti-sense ODN with a metal complex, *e.g.* terpyridylCu(II), capable of mediating mRNA hydrolysis are described in Bashkin *et al.* (1995) <u>Appl</u> Biochem Biotechnol 54:43-56.

GENETICALLY ALTERED CELL OR ANIMAL MODELS FOR K+HNOV FUNCTION

The subject nucleic acids can be used to generate transgenic animals or site specific gene modifications in cell lines. Transgenic animals may be made through homologous recombination, where the normal *K+Hnov* locus is altered. Alternatively, a nucleic acid construct is randomly integrated into the genome. Vectors for stable integration include plasmids, retroviruses and other animal viruses, YACs, and the like.

10

15

20

25

30

The modified cells or animals are useful in the study of K+Hnov function and regulation. For example, a series of small deletions and/or substitutions may be made in the K+Hnov gene to determine the role of different transmembrane domains in forming multimeric structures, ion channels, etc. Of interest are the use of K+Hnov to construct transgenic animal models for epilepsy and other neurological defects, where expression of K+Hnov is specifically reduced or absent. Specific constructs of interest include anti-sense K+Hnov, which will block K+Hnov expression, expression of dominant negative K+Hnov mutations, etc. One may also provide for expression of the K+Hnov gene or variants thereof in cells or tissues where it is not normally expressed or at abnormal times of development.

DNA constructs for homologous recombination will comprise at least a portion of the K+Hnov gene with the desired genetic modification, and will include regions of homology to the target locus. DNA constructs for random integration need not include regions of homology to mediate recombination. Conveniently, markers for positive and negative selection are included. Methods for generating cells having targeted gene modifications through homologous recombination are

known in the art. For various techniques for transfecting mammalian cells, see Keown et al. (1990) Methods in Enzymology 185:527-537.

For embryonic stem (ES) cells, an ES cell line may be employed, or embryonic cells may be obtained freshly from a host, e.g. mouse, rat, guinea pig, etc. Such cells are grown on an appropriate fibroblast-feeder layer or grown in the presence of leukemia inhibiting factor (LIF). When ES or embryonic cells have been transformed, they may be used to produce transgenic animals. After transformation, the cells are plated onto a feeder layer in an appropriate medium. Cells containing the construct may be detected by employing a selective medium. After sufficient time for colonies to grow, they are picked and analyzed for the occurrence of homologous recombination or integration of the construct. Those colonies that are positive may then be used for embryo manipulation and blastocyst injection. Blastocysts are obtained from 4 to 6 week old superovulated females. The ES cells are trypsinized, and the modified cells are injected into the blastocoel of the blastocyst. After injection, the blastocysts are returned to each uterine horn of pseudopregnant females. Females are then allowed to go to term and the resulting offspring screened for the construct. By providing for a different phenotype of the blastocyst and the genetically modified cells, chimeric progeny can be readily detected.

The chimeric animals are screened for the presence of the modified gene and males and females having the modification are mated to produce homozygous progeny. If the gene alterations cause lethality at some point in development, tissues or organs can be maintained as allogeneic or congenic grafts or transplants, or in *in vitro* culture. The transgenic animals may be any non-human mammal, such as laboratory animals, domestic animals, etc. The transgenic animals may be used in functional studies, drug screening, *etc.*, *e.g.* to determine the effect of a candidate drug on Ras or related gene activation, oncogenesis, *etc.*

20

TESTING OF K+HNOV FUNCTION and RESPONSES

Potassium channels such as K+Hnov polypeptides are involved in multiple biologically important processes. Pharmacological agents designed to affect only specific channel subtypes are of particular interest. Presently available compounds tend to be non-specific and elicit both positive and negative responses, thereby reducing clinical efficacy.

The subject polypeptides may be used in *in vitro* and *in vivo* models to test the specificity of novel compounds, and of analogs and derivatives of compounds known to act on potassium channels. Numerous pharmacological agents have profound affects on K+ channel activity. As examples, Sotalol (BETAPACE) is a class III antiarrhythmic drug that prolongs cardiac action potentials by inhibiting delayed rectifier K+ channels. Sulfonylurea drugs, such as Glipizide (GLUCOTROL) and Tolazamide (TOLAMIDE) function as antidiabetic drugs by blocking ATP-sensitive K+ channels present in pancreatic islet cells, thereby regulating insulin secretion. Diazoxide (HYPERSTAT IV) is an antihypertensive drug that activates ATP-sensitive K+ channels, resulting in the relaxation of vascular smooth muscle. There are several other examples of drugs that have antidiabetic, antihypertensive, or antiarrhythmic activities. A number of drugs that activate K+ channels that have been proposed as coronary vasodilators for the treatment of both vasospastic and chronic stable angina.

The availability of multiple K+ channel subunits allows *in vitro* reconstruction of functional channels, which may comprise different alpha and beta subunits. The individual components may be modified by sequence deletion, substitution, *etc.* to determine the functional role of specific domains.

20

25

30

Drug screening may be performed using an *in vitro* model, a genetically altered cell or animal, or purified K+Hnov protein, either as monomers, homomultimers or hetermultimers. One can identify ligands or substrates that bind to, modulate or mimic the action of K+Hnov. Drug screening identifies agents that provide a replacement for K+Hnov function in abnormal cells. Of particular interest are screening assays for agents that have a low toxicity for human cells. A wide variety of assays may be used for this purpose, including

monitoring cellular excitation and conductance, labeled *in vitro* protein-protein binding assays, electrophoretic mobility shift assays, immunoassays for protein binding, and the like. The purified protein may also be used for determination of three-dimensional crystal structure, which can be used for modeling intermolecular interactions.

The term "agent" as used herein describes any molecule, *e.g.* protein or pharmaceutical, with the capability of altering or mimicking the physiological function of *K+Hnov* polypeptide. Generally a plurality of assay mixtures are run in parallel with different agent concentrations to obtain a differential response to the various concentrations. Typically, one of these concentrations serves as a negative control, *i.e.* at zero concentration or below the level of detection.

Candidate agents encompass numerous chemical classes, though typically they are organic molecules, preferably small organic compounds having a molecular weight of more than 50 and less than about 2,500 daltons. Candidate agents comprise functional groups necessary for structural interaction with proteins, particularly hydrogen bonding, and typically include at least an amine, carbonyl, hydroxyl or carboxyl group, preferably at least two of the functional chemical groups. The candidate agents often comprise cyclical carbon or heterocyclic structures and/or aromatic or polyaromatic structures substituted with one or more of the above functional groups. Candidate agents are also found among biomolecules including peptides, saccharides, fatty acids, steroids, purines, pyrimidines, derivatives, structural analogs or combinations thereof.

Candidate agents are obtained from a wide variety of sources including libraries of synthetic or natural compounds. For example, numerous means are available for random and directed synthesis of a wide variety of organic compounds and biomolecules, including expression of randomized oligonucleotides and oligopeptides. Alternatively, libraries of natural compounds in the form of bacterial, fungal, plant and animal extracts are available or readily produced. Additionally, natural or synthetically produced libraries and compounds are readily modified through conventional chemical, physical and biochemical mans, and may be used to produce combinatorial libraries. Known

pharmacological agents may be subjected to directed or random chemical modifications, such as acylation, alkylation, esterification, amidification, etc. to produce structural analogs.

Where the screening assay is a binding assay, one or more of the molecules may be joined to a label, where the label can directly or indirectly provide a detectable signal. Various labels include radioisotopes, fluorescers, chemiluminescers, enzymes, specific binding molecules, particles, e.g. magnetic particles, and the like. Specific binding molecules include pairs, such as biotin and streptavidin, digoxin and antidigoxin etc. For the specific binding members, the complementary member would normally be labeled with a molecule that provides for detection, in accordance with known procedures.

10

15

20

25

30

A variety of other reagents may be included in the screening assay. These include reagents like salts, neutral proteins, e.g. albumin, detergents, etc that are used to facilitate optimal protein-protein binding and/or reduce non-specific or background interactions. Reagents that improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc. may be used. The mixture of components are added in any order that provides for the requisite binding. Incubations are performed at any suitable temperature, typically between 4 and 40°C. Incubation periods are selected for optimum activity, but may also be optimized to facilitate rapid high-throughput screening. Typically between 0.1 and 1 hours will be sufficient.

The compounds having the desired pharmacological activity may be administered in a physiologically acceptable carrier to a host in a variety of ways, orally, topically, parenterally e.g. subcutaneously, intraperitoneally, by viral infection, intravascularly, etc. Depending upon the manner of introduction, the compounds may be formulated in a variety of ways. The concentration of therapeutically active compound in the formulation may vary from about 0.1-100 wt.%. The pharmaceutical compositions can be prepared in various forms, such as granules, tablets, pills, suppositories, capsules, suspensions, salves, lotions and the lik. Pharmaceutical grade organic or inorganic carriers and/or diluents suitable for oral and topical use can be used to make up

compositions containing the therapeutically-active compounds. Diluents known to the art include aqueous media, vegetable and animal oils and fats. Stabilizing agents, wetting and emulsifying agents, salts for varying the osmotic pressure or buffers for securing an adequate pH value, and skin penetration enhancers can be used as auxiliary agents.

5

10

15

20

25

30

It is to be understood that this invention is not limited to the particular methodology, protocols, cell lines, animal species or genera, and reagents described, as such may vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments only, and is not intended to limit the scope of the present invention which will be limited only by the appended claims.

As used herein the singular forms "a", "and", and "the" include plural referents unless the context clearly dictates otherwise. Thus, for example, reference to "a cell" includes a plurality of such cells and reference to "the cell" includes reference to one or more cells and equivalents thereof known to those skilled in the art, and so forth. All technical and scientific terms used herein have the same meaning as commonly understood to one of ordinary skill in the art to which this invention belongs unless clearly indicated otherwise.

It must be noted that as used herein and in the appended claims, the singular forms "a", "and", and "the" include plural referents unless the context clearly dictates otherwise. Thus, for example, reference to "a complex" includes a plurality of such complexes and reference to "the formulation" includes reference to one or more formulations and equivalents thereof known to those skilled in the art, and so forth.

All publications mentioned herein are incorporated herein by reference for the purpose of describing and disclosing, for example, the methods and methodologies that are described in the publications which might be used in connection with the presently described invention. The publications discussed above and throughout the text are provided solely for their disclosure prior to the filing date of the present application. Nothing herein is to b construed as an

admission that the inventors are not entitled to antedate such disclosure by virtue of prior invention.

EXPERIMENTAL

The following examples are put forth so as to provide those of ordinary skill in the art with a complete disclosure and description of how to make and use the subject invention, and are not intended to limit the scope of what is regarded as the invention. Efforts have been made to ensure accuracy with respect to the numbers used (e.g. amounts, temperature, concentrations, etc.) but some experimental errors and deviations should be allowed for. Unless otherwise indicated, parts are parts by weight, molecular weight is average molecular weight, temperature is in degrees centigrade; and pressure is at or near atmospheric.

15 Methods

5

Two different types of sequence searches were performed. The first centered on the most highly conserved region of the K+ channel family, the pore domain. The pore is composed of 15-17 amino acids and can be divided into subfamilies based on the number of transmembrane segments present in the channel. Eleven variant peptide sequences corresponding to the pore domain were used in TBLASTN searches against the EST division of Genbank. Significant matches were identified, and classified into 2 categories: identical to known human K+ channels and related to known K+ channels. The pore sequences are shown in Table 2.

TABLE 2

	_	
SEG ID NO	Gendank #	
49	L02751	TGGTGGGCTGTGGTGACCATGACAACTGTGGGCTATGGGGACATG
50	M60451	TGGTGGGCAGTGGTCACCATGACCACTGTGGGCTACGGGGACATG
51	L02752	TGGTGGGCAGTCGTCTCCATGACAACTGTAGGCTATGGAGACATG
52	M55515	TGGTGGGCAGTGGTAACCATGACAGTGGGGTTACGGCGATATG
53	211585	TGGTGGGCTGTGGTCACCATGACGACCCTGGGGCTATGGAGACATG
54	U40990	TGGTGGGGGTGGTCACAGTCACCATCGGCTATGGGGACAAG
55	126643	TOGTGGGCAGTGGTCATGACCACGGTTGGCTATGGGGACATG
58	M96747	TGGTGGGCCGTGGTCACCATGACGACCCTGGGGCTATGGAGACATG
57	M64676	TOGTGGGCTGTGGTCACCATGACGACACTGGGCTACGGAGACATG
58	M55514	TGGTGGGCTGTGGTGACCATGACCACTGTGGGGCTATGGGGGACATG
59	X83582	TTCCTGTTCCCATTGAGACCGAAACAACCATTGGGTATGGCTTCCG
80	S78684	TTTTTATTCTCAATAGAGACAGAAACCACCATTGGTTATGGCTACCG
81	U22413	TTCCTCTTCTCCATTGAGACCCAGACCATAGGCTATGGTTTCAG
62	U24056	TTCCTGTTCCGGTGGAGGCGAGGGACCATCGGCTATGGGTTCCG
83	U52155	TTCCTCTTCCCTTGAATCCCAAACCACCATTGGCTATGGCTTCCG
2	D87291	TITCTCTTTTCCCTGGAATCCCAGACAACCATTGGCTATGGAGTCCG
85	D50582	TTCCTTTTCTCCATTGAGGTCCAAGTGACTATTGGCTTTGGGGGGCG
88	D50315	TTTCTCTTCTCCATTGAAGTTCAAGTTACCATTGGGTTTGGAGGGAG
67	U04270	GCGCTCTACTTCACCTTCAGCAGCCTCACCAGTGTGGGCTTCGGCAAC

The unique pore peptides sequences are shown in Table 3.

TABLE 3

and the same of							
SEQ ID NO	Amino acid sequence						
68	WWAVVSMTTVGYGDM						
69 WWAVVTMTTLGYGDM 70 WWGWTVTTIGYGDK							
70	WWGVVTVTTIGYGDK						
71	WWAVVTMTTVGYGDM						
72	FLFSIEVQVTIGFGG						
73	FLFSLESQTTIGYGV						
74	FLFSIETETTIGYGY						
75	FLFSIETQTTIGYGF						
76	FLFSVETQTTIGYGF						
77	FLFSLESQTTIGYGF						
78 FLFSIETETTIGYGF							
79	ALYFTFSSLTSVGFGN						

The second set of experiments was based on a complex, reiterative process.

Annotated protein and DNA sequences were obtained from GenBank for all known K+ channels from all species. The TBLASTN and BLASTN programs were used to identify homologous ESTs, which were then analyzed using the BLASTX and BLASTN algorithms to identify ESTs which were related to K+ channels yet not identical to any known human K+ channel gene.

Novel human K+ channels were defined as those that had clear homology to known K+ channels from any species and were not present as identities or near identities to any human-derived sequences in any division of Genbank.

Isolation of full length cDNA sequence. EST clones were picked from the IMAGE consortium cDNA library and end-sequenced with vector primers. Gap closure was achieved either by primer walking or transposon sequencing. GeneTrapper (Life

15

Technologies) was used to isolate larger cDNA clones according to the provided protocol. RACE was used to extend the sequences as necessary using standard protocols.

Sequences were assembled in Sequencher (Gene Codes). The presence of open reading frames was assessed as well as potential start codons. Potential polymorphisms were detected as sequence variants between multiple independent clones. Sequence homologies were detected using the BLAST algorithms.

The completed gene sequences and predicted amino acid sequences are provided as SEQ ID NO:1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21-24, 26 and 28-29. Polymorphisms, chromosome locations and family assignments are shown in Table 1.

10

ESTs that had top human hits with >95% identity over 100 amino acids were discarded. This was based upon the inventors' experience that these sequences were usually identical to the starting probe sequences, with the differences due to sequence error. The remaining BLASTN and BLASTX outputs for each EST were examined manually, i.e., ESTs were removed from the analysis if the inventors determined that the variation from the known related probe sequence was a result of poor database sequence. Poor database sequence was usually identified as a number of 'N' nucleotides in the database sequence for a BLASTN search and as a base deletion or insertion in the database sequence, resulting in a peptide frameshift, for a BLASTX output. ESTs for which the highest scoring match was to non-related sequences were also discarded at this stage. The EST sequences that correspond to each clone are shown in Table 4.

Table 4

Genbank Accession#	K+Hnov	clone ID	Trace	IMAGE Plate Coordinates	Read 5'/3'
N39619	K+Hnov2	277113	yy51h05.s1	611p10	3'
N46767	K+Hnov2	277113	yy51h05.r1	611p10	5'
R19352	K+Hnov11	33144	yg24f12.r1	155024	5'
R44628	K+Hnov11	33144	yg24f12.s1	155024	3'

R35526	K+Hnov14	37299	yg64e08.r1	165015	5'
R73353	K+Hnov14	157854	yl10e04.r1	251g07	5'
AA397616	K+Hnov14	728558	zt79c08.r1	1787j15	5'
AA286692	K+Hnov28	700757	zs48h03.r1	1715d6	5'
AA150494	K+Hnov42	491748	zl08e07.s1	1170013	3'
AA156697	K+Hnov42	491748	zl08e07.r1	1170013	5'
AA191752	K+Hnov42	626699	zp82d06.r1	1522f12	5'
AA216446	K+Hnov42	626699	zp82d06.s1	1522f12	3'
AA430591	K+Hnov42	773611	zw51f10.r1	1904020	5'
AA236930	K+Hnov44	683888	zs01a05.s1	1671e9	3'
AA236968	K+Hnov44	683888	zs01a05.r1	1671e9	5'

EXAMPLE 2: CHROMOSOMAL LOCALIZATION

Two primers were designed in the 3'-untranslated regions of each gene sequence to amplify a product across the Stanford G3 radiation hybrid map, or the Whitehead GB4 panel. The PCR data were submitted for automatic two-point analysis. Mapping data were correlated with cytoband information and comparisons with the OMIM human gene map data base were made. The following primers were made:

K+Hnov1 on GB4

(SEQ ID NO:31) F: 5' TATCCACATCAATGGACAAAGC 3'
(SEQ ID NO:32) R: 5' TGCATAACTGGCTGGGTGTA 3'
Results: 1.71 cR from D2S331, Cytogenetic location of 2q37

K+Hnov2 on G3

15 F: 5' GTCAGGTGACCGAGTTCA 3'
R: 5' GCTCCATCTCCAGATTCTTC 3'
Results: 0.0 cR from SHGC-1320, Cytogenetic location of 11q12

Results: 0.0 cR from SHGC-1320, Cytogenetic location or 114.

K+Hnov6 on GB4
(SEQ ID NO:33) F: 5' TGACATCACTGGATGAACTTGA 3'
(SEQ ID NO:34) R: 5' TGCCTGCAAAGTTTGAACAT 3'
Results: 5.23 cR from WI-5509, Cytogenetic location of 2p23

K+Hnov9 on GB4
(SEQ ID NO:35) F: 5' TGACATCACTGGATGAACTTGA 3'
(SEQ ID NO:36) R: 5' TGCCTGCAAAGTTTGAACAT 3'

Results: 1.21 cR from AFM200VC7, Cytogenetic location of 8q23

K+Hnov11 on GB4

(SEQ ID NO:37) F: 5' ACCTGGTGGTATGGAAGCAT 3' (SEQ ID NO:38) R: 5' TTTCTCCTGGCCTCTACCC 3'

Results: 2.43 cR from WI-6756, Cytogenetic location of 8q23

K+Hnov12 on G3

(SEQ ID NO:39) F: 5' TCCCTCTTGGGTGACCTTC 3'

10 (SEQ ID NO:40) R: 5' ATCTTTGTCAGCCACCAGCT 3'

Results: 7.45 cR from SHGC-32925, Cytogenetic location of Xp21

K+Hnov14 on GB4

(SEQ ID NO:41) F: 5' AGGTGTGCTGCCATCTGCTGTTCG3'

15 (SEQ ID NO:42) R: 5' AGCCTATCCTCTGAGAGTCAGG

Results: 7.69 cR from WI-7107, Cytogenetic location of 12q14

K+Hnov28 on GB4

(SEQ ID NO:43) F: 5' AAGCAGAGTACTCATGATGCC 3'

20 (SEQ ID NO:44) R: 5' TCTGGTAGACAGTACAGTGG 3'

Results: 35.38 cR from WI-9695, Cytogenetic location of 3q29

K+Hnov42 on G3

(SEQ ID NO:45) F: 5' CATTTGGCTGGTCCAAGATG 3'

25 (SEQ ID NO:46) R: 5' AGTCATTGGTAGGGAGGTAC 3'

Results: 7.45 cR from SHGC-32925, Cytogenetic location of Xp21

K+Hnov44 on G3

(SEQ ID NO:47) F: 5' CATGCTTCTACAGTCCAGCC 3'

30 (SEQ ID NO:48) R: 5' GGTCCTCAGTTGCAGAAATC 3'

Results: 7.45 cR from SHGC-32925, Cytogenetic location of Xp21

Map positions for K+Hnov15 and K+Hnov27 were obtained from public databases. K+Hnov2 and K+Hnov4 have not been mapped.

35

EXAMPLE 3: EXPRESSION ANALYSIS

RT-PCR was utilized to characterize the expression pattern of the novel ion channels. This approach used RNA from 30 different tissues to generate first strand cDNA. Total RNA was purchased (Clontech, Invitrogen) and used to synthesize first strand cDNA using M-MLV reverse transcriptase and the supplied buffer (Gibco-BRL). The 20 µl reaction contained 5 µg total RNA, 100 ng of random primers, 10 mM DTT.

0.5 mM each dNTP, and an RNAse inhibitor (Gibco-BRL). Identical reactions were set up without reverse transcriptase to control for DNA contamination in the RNA samples. The synthesis reaction proceeded for 1 hour at 37°C followed by 10 minutes at 95°C. These cDNAs, along with control cDNA synthesis reactions without reverse transcriptase, were diluted 1:5 and 2 µl of each sample were arrayed into 96-well trays, dried, and resuspended in PCR buffer prior to PCR amplification. The cDNAs were tested with primers with defined expression patterns to verify the presence of amplifiable cDNA from each tissue. Gene-specific primers were used to amplify the cDNAs in 20 µl PCR reactions with standard conditions, 2.5 mM MgCl₂, Taq Gold, and an appropriate annealing temperature.

This approach provides for relatively high-throughput analysis of gene expression in a large set of tissues in a cost-efficient manner and provides qualitative analysis of gene expression only. Modifications can be employed, such as the use of internal control primers, limited cycling parameters, and dilution series to convert this to a quantitative experiment.

10

Table 3

WO 99/43696

terus	•	•	•	•		٠	•	٠	٠	٠	٠		I	•		
Inschea	٠	•	•	٠	•	٠	\cdot	٠	٠	•	٠	Ī		•		
Thymus	•	٠		•	٠	•	•	٠	•	٠	٠	Ī	I	-		
Testis	•	•		•	٠	Γ		٠	٠	•	٠	T	Ī	7		
Stomach	•	1		٠	ŀ	٠	1	•	٠	٠	٠	T		•		sample
Spleen	•		1	•	٠		1	1	٠	٠	1	1	1	•		Sa
Smail	٠	1.	•	•	•	1		•	٠	٠	1	·		٠		it i
Intestine Skin	r	T.	T	1.		T	1.	•	T	1	T	Ī		٠		Ş
Skeletul	١,	†	1.	1.	t	t	t.	1.	1.	1.	†	,		•		T
Muscle	Ľ	Ľ	L	L	L	L	1	╄	╄	+	4	4	4	┝	ł	9
Salivary	١,	Τ,	1.		1.		١.	1.	١.	١,	١,	٠l		+	1	
Gland	ľ	Τ.	T	Ľ	Ľ	Ľ	L	L	L	L	L	4	_	L	1	ě
Rectum	Γ	1.	Ī	1.	1	T	1	1	·T	1	·			٠		100
Prostat-	ļ	, ,	٠,	.†•	†	1.	.†.	1.				٠		٠		ted to date on acted per and
Placent :	ŀ	,	+	. .	1	ŀ	•	ŀ	·I·	·ŀ	·	+		ŀ		
Pancreas	ŀ	·Ī	$\overline{\cdot}$		·I·	•]•	·	·ŀ	$\cdot \cdot$	•	•	+	L	ŀ		1
Mammary Gland		•	•	• [·T	•	•	.[•	٠	٠	+		ŀ	1	4 1.1.
Lung	ı	•	•				•	·T	•	•	٠	+		ŀ	-	1
Liver		•	+	•	·	•	•	•	•	·	٠	*		ŀ	[
Ridne+		•	-	-	•	•	•	\cdot	٠	٠	٠	٠		ŀ		
HeLa C		•	٠	·	\cdot	•	٠	\cdot	٠	٠	+	ŀ		ŀ	1	
Heart		٠	٠	\cdot	+	•	٠	\cdot	·	٠	٠	ŀ	1	ŀ	1	
Fetal c ↔ 1		٠	٠	\cdot	\cdot	•	٠	·	+	٠	+	ŀ		ŀ	1	
Fetal Bri		٠	•	+	٠	•	٠	٠	٠	٠	÷	ŀ	1	1	1	
Escondi		٠	+		+			·	٠		·	ŀ	ŀ		1	
Colon		٠	٠	٠	+	•	•	•	٠	Ŀ	٠	Ŀ	ŀ	_	<u>.</u>	
Cer		•	٠					•	٠	L	٠	1			1	
Cerebo		٠	٠	٠	+	٠	٠	·	ŀ	•	٠	ŀ	·		Ċ	
Brain		٠	٠	٠	٠	·	٠	ŀ	ŀ	•	ŀ	ŀ	٠		•	
Bladder		٠	٠	ŀ	·	ŀ	•	٠	•	•	ŀ	ŀ	٠		٠	
Adrerat		Г	Τ	١.	1	1			1.	. .	١,	١.	۱	ľ	٠	
Gland		•	1	ľ	١	ľ		Ľ	Ĺ	Ľ	L	J	ا	L	┖	1
Actocse		ļ.	1.	T	T	T	T	1		1	T.	•			٠	
Ancher nar	٠,.		t	t	t	1	1	T	T	1	1		j		Γ	
	-	1.000	COLLAN	A.Hook	A 1400 A		No. House	2		No. Union	CI AGULLA	K+Hnov2/	K+Hnov28	K+Hnov42	K+Hnov44	

A *+" indicates expression in the tissue, a *-" indicates no expression, and blank square indicates no data for that sample.

K+Hnov49 on Whitehead GB4 RH mapping panel:

Primer 1 (SEQ ID NO:5): 5' - CATAGCCATAGGTGAGGACT - 3'

Primer 2: (SEQ ID N:6) 5' - GAGAGGAAAACAGTCTGGGC - 3'

Results: Cytogenetic location 1q41, 4.6cR from framework marker D1S217

K+Hnov59 on Whitehead GB4 RH mapping panel

Primer 1 (SEQ ID NO:7): 5' - GGACATCGAACTAAGACCTG - 3'

Primer 2 (SEQ ID NO:8): 5' - TCCCATGCCATTCAGATCTG - 3'

10 Results: Cytogenetic location 19q13.2, 8.34cr from framework marker D19S425

EXPRESSION ANALYSIS OF K+HNOV49

20

A probe was created from a fragment corresponding to nucleotides 50 to 1284 of SEQ ID NO:83 (K+Hnov49) and purified DNA fragment was labeled with [³²P]dCTP (Amersham) by the random primer method. Adult human Multiple Tissue Northern (MTM™) Blots (Clontech) were hybridized with the [³²P]-labeled fragment in ExpressHyb™ solution (Clontech) for four hours, washed to a final stringency of 0.1xSSC, 0.1% SDS at 65°C and subjected to autoradiography for 24 hours.

Analysis revealed that K+Hnov49 is expressed as an approximately 4.2kb mRNA. Expression levels of K+Hnov49 are high in brain and liver and low in kidney tissues. No mRNA was detectable on these Northern blots for heart, skeletal muscle, colon, thymus, spleen, small intestine, placenta, lung or peripheral blood leukocytes indicating either a very low level of expression or that it is not expressed in these tissues. Expression analysis was also carried out by RT-PCR across an extended series of tissues. The results of these analyses are shown in Table 4. Primer pairs used for amplification of K+Hnov49 and 59 are the same as those used for RH mapping as indicated above.

Table 4

	Adipose	Adrenal Gland	Bladder	Brain	Cerebellum	Cervix	Colon	Esophagus	Fetal Brain	Fetal Liver	Heart	He La Cell	Kidney	Liver	Lung	Mammary Gland	Pancreas	Placenta	Prostate	Rectum	Salivary Gland	Skeletal Muscle	Skin	Small Intestine	Spleen	Stomach	Testus		Trachea	
#49	+	+	+	+	+	+	-	+	+		+	+	+	-	+	+	-	-	+	-	+	+	-	+	-	+	+	+	-	-
#59	_	_				+	_	+		+	+			+	+	+	+	_	+	+	+	_	_	+	. +	. +	+	+	+ -	+

WHAT IS CLAIMED IS:

- 1. An isolated nucleic acid encoding a mammalian K+Hnov protein.
- 2. An isolated nucleic acid according to Claim 1, wherein said K+Hnov protein has the amino acid sequence of SEQ ID NO:2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 25, 27, 30, 81 or 83.
 - 3. An isolated nucleic acid according to Claim 1, wherein said K+Hnov protein has an amino acid sequence that is substantially identical to the amino acid sequence of SEQ ID NO:2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 25, 27, 30, 81 or 83.
 - 4. An isolated nucleic acid according to Claim 1 wherein the nucleotide sequence of said nucleic acid is SEQ ID NO:1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 22, 23, 24, 26, 28, 29, 80 or 82.
 - 5. An isolated nucleic acid that hybridizes under stringent conditions to a nucleic acid sequence of claim 4.
- 20 6. An expression cassette comprising a transcriptional initiation region functional in an expression host, a nucleic acid having a sequence of the isolated nucleic acid according to Claim 1 under the transcriptional regulation of said transcriptional initiation region, and a transcriptional termination region functional in said expression host.

25

10

15

7. A cell comprising an expression cassette according to Claim 6 as part of an extrachromosomal element or integrated into the genome of a host cell as a result of introduction of said expression cassette into said host cell, and the cellular progeny of said host cell.

8. A method for producing mammalian K+Hnov protein, said method comprising:

growing a cell according to Claim 7, whereby said mammalian K+Hnov protein is expressed; and

- isolating said K+Hnov protein free of other proteins.
- 9. A purified polypeptide composition comprising at least 50 weight % of the protein present as a K+Hnov protein or a fragment thereof.
- 10. A monoclonal antibody binding specifically to a K+Hnov protein.
 - 11. A non-human transgenic animal model for K+Hnov gene function wherein said transgenic animal comprises an introduced alteration in a K+Hnov gene.

15

- 12. The animal model of claim 11, wherein said animal is heterozygous for said introduced alteration.
- 13. The animal model of claim 12, wherein said animal is homozygous for said introduced alteration.
 - 14. The animal model of claim 12, wherein said introduced alteration is a knockout of endogenous K+Hnov gene expression.

SEQUENCE LISTING

<110> Miller, Andrew Curran, Mark Buckler, Alan	
<120> Novel Human Potassium Channels	
<130> SEQ-15PCT	
<150> 60/076,687 <151> 1998-02-25	
<150> 60/095,836 <151> 1998-08-07	
<150> 60/116,448 <151> 1999-01-19	
<160> 87	
<170> FastSEQ for Windows Version 3.0	
<210> 1 <211> 2932 <212> DNA <213> H. sapiens	
<220> <221> CDS <222> (103)(1180) <223> K+Hnov1	
<pre><400> 1 attaaaatta tctgatcaaa aaggcagact ctgtaaattt ccttaagacc taccttggca attaaaaggctga cccagcaaaa gaactgagaa atacagcctg ag atg gac agc agt</pre>	60 114
aat tgc aaa gtt att gct cct ctc cta agt caa aga tac cgg agg atg Asn Cys Lys Val Ile Ala Pro Leu Leu Ser Gln Arg Tyr Arg Arg Met 5 10 15 20	162
gtc acc aag gat ggc cac agc aca ctt caa atg gat ggc gct caa aga Val Thr Lys Asp Gly His Ser Thr Leu Gln Met Asp Gly Ala Gln Arg 25 30 35	210
ggt ctt gca tat ctt cga gat gct tgg gga atc cta atg gac atg cgc Gly Leu Ala Tyr Leu Arg Asp Ala Trp Gly Ile Leu Met Asp Met Arg 40 45 50	258
tgg cgt tgg atg atg ttg gtc ttt tct gct tct ttt gtt gtc cac tgg Trp Arg Trp Met Met Leu Val Phe Ser Ala Ser Phe Val Val His Trp 55 60 65	306
ctt gtc ttt gca gtg ctc tgg tat gtt ctg gct gag atg aat ggt gat Leu Val Phe Ala Val Leu Trp Tyr Val Leu Ala Glu Met Asn Gly Asp 70 75 80	354

ctg gaa cta gat cat gat gcc cca cct gaa aac cac act atc tgt gtc Leu Glu Leu Asp His Asp Ala Pro Pro Glu Asn His Thr Ile Cys Val 85 90 95 100	402
aag tat atc acc agt ttc aca gct gca ttc tcc ttc tcc ctg gag aca Lys Tyr Ile Thr Ser Phe Thr Ala Ala Phe Ser Phe Ser Leu Glu Thr 105 110 115	450
caa ctc aca att ggt tat ggt acc atg ttc ccc agt ggt gac tgt cca Gln Leu Thr Ile Gly Tyr Gly Thr Met Phe Pro Ser Gly Asp Cys Pro 120 125 130	498
agt gca atc gcc tta ctt gcc ata caa atg ctc cta ggc ctc atg cta Ser Ala Ile Ala Leu Leu Ala Ile Gln Met Leu Leu Gly Leu Met Leu 135 140 145	546
gag gct ttt atc aca ggt gct ttt gtg gcg aag att gcc cgg cca aaa Glu Ala Phe Ile Thr Gly Ala Phe Val Ala Lys Ile Ala Arg Pro Lys 150 155 160	594
aat cga gct ttt tca att cgc ttt act gac aca gca gta gta gct cac Asn Arg Ala Phe Ser Ile Arg Phe Thr Asp Thr Ala Val Val Ala His 165 170 175 180	642
atg gat ggc aaa cct aat ctt atc ttc caa gtg gcc aac acc cga cct Met Asp Gly Lys Pro Asn Leu Ile Phe Gln Val Ala Asn Thr Arg Pro 185 190 195	690
agc cct cta acc agt gtc cgg gtc tca gct gta ctc tat cag gaa aga Ser Pro Leu Thr Ser Val Arg Val Ser Ala Val Leu Tyr Gln Glu Arg 200 205 210	738
gaa aat ggc aaa ctc tac cag acc agt gtg gat ttc cac ctt gat ggc Glu Asn Gly Lys Leu Tyr Gln Thr Ser Val Asp Phe His Leu Asp Gly 215 220 225	786
atc agt tct gac gaa tgt cca ttc ttc atc ttt cca cta acg tac tat Ile Ser Ser Asp Glu Cys Pro Phe Phe Ile Phe Pro Leu Thr Tyr Tyr 230 235 240	834
cac tcc att aca cca tca agt cct ctg gct act ctg ctc cag cat gaa His Ser Ile Thr Pro Ser Ser Pro Leu Ala Thr Leu Leu Gln His Glu 245 250 255 260	882
aat cct tct cac ttt gaa tta gtt gta ttc ctt tca gca atg cag gag Asn Pro Ser His Phe Glu Leu Val Val Phe Leu Ser Ala Met Gln Glu 265 270 275	930
ggc act gga gaa ata tgc caa agg agg aca tcc tac cta ccg tct gaa Gly Thr Gly Glu Ile Cys Gln Arg Arg Thr Ser Tyr Leu Pro Ser Glu 280 285 290	978
atc atg tta cat cac tgt ttt gca tct ctg ttg acc cga ggt tcc aaa Ile Met Leu His His Cys Phe Ala Ser Leu Leu Thr Arg Gly Ser Lys 295 300 305	1026
ggt gaa tat caa atc aag atg gag aat ttt gac aag act gtc cct gaa Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Thr Val Pro Glu 310 315 320	1074
ttt cca act cct ctg gtt tct aaa agc cca aac agg act gac ctg gat	1122

Phe Pro Thr Pro Leu Val Ser Lys Ser Pro Asn Arg Thr Asp Leu Asp	
atc cac atc aat gga caa agc att gac aat ttt cag atc tct gaa act Ile His Ile Asn Gly Gln Ser Ile Asp Asn Phe Gln Ile Ser Glu Thr	1170
345	1220
gga ctg aca g aataagactt atccattttt taatgtatta aatacaccca Gly Leu Thr	
gccagttatg cagctacttt ttctttactg tatctcatgt tttcttttt caatgctaat	1280 1340
gecagttatg cagetacttt ttetttaetge tateedatge teasgaatgge tgagetaaca tatagetete tacateaegg taateatgee tatgeetaca taagaatgge tgagataaca	1400
tatagetete tacateaegg taateatgee tatgeteata tadgatetete tagaateaat atacaacatte tagaaacata acaetetaca ttacaaagat tagtaacttat aagaatgtag	1460
gtaactcaac ttgacagaca cttatacaga aatgacagttt aacacaactg aactctaaga	1520
tatgatacta gtaatgaagg caaaatggac aggatta agtttcaaac	1580
agatcacca ttaatctctc attituates betagt gastatcctt taagagctgg	1640
ctageteect gggtggaatg acgaetteat tatactatga aacttgagta	1700 1760
gattttttte aagacaacaa agatcattca tttggttttt tatacaagagaaa aacaacaag agtattacct ccttaatttt taacaactaa gaacaaaaat taacgagaaa aacaacaag agtattacct ccttaatttt taacaactaa gaacaacaa atacatatat	1820
agtattacct ccttaatttt taacaactaa gaacaadaat taacgagatta tacatatat tacagattta tacataaacc taaaagcatt tgaacatgac acccgaacac atacatatat tacagattta tacataaacc aaaagcatta ataagctcca gcccaaatgg aacctgtggg	1880
tacagattta tacataaacc taaaagcatt tgaacatgat accegatata gttcacttat ttgtggcaga aggtgatcag ataagctcca gcccaaatgg aacctgtggg	1940
gttcacttat ttgtggcaga aggtgatcag ataageteca gtecaadass segurate aggtgtatttt gcattgcaag gagacgcaaa attttatttt	2000
tcaaacqqtq attcatctaa atgattteta santatata aattcataaa tacaaaaata	2060
gtatgattca titgatcata tadadacatta garagata actroagtga gtagggaaaa	2120
tatttaagtc tttatagata taaagettta taaagettta agcattttca aactcaaatt	2180
aaatctacag tagataaagc aaaagataat taggcaataa agcattatacacagt cctgtttcca acttcaaata gttttttcta taaacacaaa atcagtgttt attcaccagt	2240 2300
cctgtttcca acttcaaata gttttttcta taadatataa atcagtc tataaaaatt aggaggttgg actagatgaa ctctattatt tctttctaaa tctaatagtc tataaaaatt aggaggttgg actagatgaa ctctattatt ctaatagag ccctttccct tatgtccagt	2360
aggaggttgg actagatgaa ctctattatt tetttetaa tetateget tatgteeagt atgtteete tgtttttat tttatetatg etaaaatgag ecettteeet tatgteeagt atgttteete tgtttttat tetatetatg etaaaatgag aaggaaact gteettaaaa	2420
atgittcctc tgttttttat tttatctatg ctaadatgag cccertagaaa ttaagatgat catttgcatg attttcattt caataaaaaa aagagaaact gtccttaaaa ttaagatgat catttgcatg attttcattt caataaaaaa caqatttgtg gctgttcttt	2480
ttaagatgat catttgcatg attttcattt caatadada augustttgtg gctgttcttt caaaacaaaa accaaaaaag tcaccctatc aggtttcaaa cagatttatg gctgttcttt caaaacaaaa	2540
totgaaattt coottattoa ggittotgeg sattantagt agotgagtaa aggcaagcag	2600
gtgatgacct aggcaggaat catctctty data the transcript chattracc aaaaggaaaa	2660
gtgtgaagag cagggctcag cagcaagtca catttttcta ttactogata gtgtgaagag cagggctcag gagtggtcta agactgataa tagcagaaga atatcaagaa gaaaataaag aagaactctg gagtggtcta tgtttgaagaa tcttagacat tcattcttaa	2720
gaaaataaag aagaactetg gagtggteta agaetgataa tegtagacat teattettaa cacagaaact taattattgt gaacttttge tgttgaaaa tettagacat teattettaa cacagaacat taattattgt gaacttttge cacagactat tgtaacacat aaagacagca	2780 2840
cacagaaact taattattgt gaacttttgc tgtttgaaaa teetagaaac aaagacagca gtagaaatca gaccaacaga ttttcccaac ccaagactat tgtaacacat aaagacagca gtagaaatca gaccaacaga gattcaccta acctttgaaa ataaagtagt	2900
	2932
attgaagact taaaaaaaaaa aaaaaaaaaa aa	
<210> 2 <211> 359	
<212> PRT	
<213> H. sapiens	
<pre><400> 2 Met Asp Ser Ser Asn Cys Lys Val Ile Ala Pro Leu Leu Ser Gln Arg 10 15</pre>	
Met Asp Ser Ser Ash Cys Lys val 113 10 15	
1 5 Tyr Arg Arg Met Val Thr Lys Asp Gly His Ser Thr Leu Gln Met Asp	
20 25 Gly Ala Gln Arg Gly Leu Ala Tyr Leu Arg Asp Ala Trp Gly Ile Leu 45	
Gly Ala Gir Alg Sar Dhe	
35 Met Asp Met Arg Trp Arg Trp Met Met Leu Val Phe Ser Ala Ser Phe 55 60	
50 55 Val Val His Trp Leu Val Phe Ala Val Leu Trp Tyr Val Leu Ala Glu Val Val His Trp Leu Val Phe Ala Val Leu Trp 75 80	
Val Val His Trp Let Val 122 75	
Met Asn Gly Asp Leu Glu Leu Asp His Asp Ala Pro Pro Glu Ash his	
85 90 95 Thr Ile Cys Val Lys Tyr Ile Thr Ser Phe Thr Ala Ala Phe Ser Phe	
105 100 105 110	

Ser Leu Glu Thr Gln Leu Thr Ile Gly Tyr Gly Thr Met Phe Pro Ser	
Gly Asp Cys Pro Ser Ala Ile Ala Leu Leu Ala Ile Gln Met Leu Leu	
Gly Leu Met Leu Glu Ala Phe Ile Thr Gly Ala Phe Val Ala Lys Ile	
145 150 155 160 145 Ala Arg Pro Lys Asn Arg Ala Phe Ser Ile Arg Phe Thr Asp Thr Ala Ala Arg Pro Lys Asn Arg Ala Phe Ser Ile Arg Phe Thr Asp Thr Ala	
Val Val Ala His Met Asp Gly Lys Pro Asn Leu Ile Phe Gin Val Ala	
Asn Thr Arg Pro Ser Pro Leu Thr Ser Val Arg Val Ser Ala Val Leu 205 207 208 209 209 200 200 200 200 200 200 200 200	
Tyr Gln Glu Arg Glu Asn Gly Lys Leu Tyr Gln Thr Ser van Asp File	
His Leu Asp Gly Ile Ser Ser Asp Glu Cys Pro Phe File 116 126	
225 230 255 Leu Thr Tyr His Ser Ile Thr Pro Ser Ser Pro Leu Ala Thr Leu	
Leu Gln His Glu Asn Pro Ser His Phe Glu Leu Val Val Phe Leu Ser 270 260 265 270 270 270	
Ala Met Gln Glu Gly Thr Gly Glu Ile Cys Gln Arg Arg Thr Ser Tyr 285 275 280 285	
Leu Pro Ser Glu Ile Met Leu His His Cys Phe Ala Ser Leu Leu III	
290 295 300 Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Ser Lys Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly Gly Glu Tyr Gln Ile Lys Met Glu Asn Phe Asp Lys Arg Gly	
Thr Val Pro Glu Phe Pro Thr Pro Leu Val Ser Lys Ser Pro Asn Arg	
Thr Asp Leu Asp Ile His Ile Asn Gly Gln Ser Ile Asp Asn Phe Gln 340 345 350	
Ile Ser Glu Thr Gly Leu Thr 355	
210. 2	
<210> 3	
<211> 1927	
<211> 1927 <212> DNA	
<211> 1927 <212> DNA <213> H. sapiens	
<211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS	
<211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908)	
<211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS	
<211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908) <223> K+Hnov4	60
<pre><211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908) <223> K+Hnov4 <400> 3 ggagccccgc agcgcttctt atgatcagct cggtgtgtgt ctcctcctac cgcgggcgca ggagccccgc agcgcttctt atgatcagct gtctgaagga ggag atg gcc aag ggc</pre>	60 116
<211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908) <223> K+Hnov4	
<pre><211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908) <223> K+Hnov4 <400> 3 ggagccccgc agcgcttctt atgatcagct cggtgtgtgt ctcctcctac cgcgggcgca ggagccccgc agcgcttctt atgatcagct gtctgaagga ggag atg gcc aag ggc</pre>	
<pre> <211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908) <223> K+Hnov4 <400> 3 ggagcccgc agcgcttctt atgatcagct cggtgtgtgt ctcctcctac cgcgggcgca agtcggggaa caagcctccg tccaaaacat gtctgaagga ggag atg gcc aag ggc Met Ala Lys Gly 1 1 1 1 1 1 1 1 1 1 1 1 1</pre>	
<pre> <211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908) <223> K+Hnov4 <400> 3 ggagccccgc agcgcttctt atgatcagct cggtgtgtgt ctcctcctac cgcgggcgca agtcggggaa caagcctccg tccaaaacat gtctgaagga ggag atg gcc aag ggc Met Ala Lys Gly 1 gag gcg tcg gag aag atc atc atc aac gtg ggc ggc acg cga cat gag Glu Ala Ser Glu Lys Ile Ile Ile Asn Val Gly Gly Thr Arg His Glu 20</pre>	116
<pre> <211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908) <223> K+Hnov4 <400> 3 ggagcccgc agcgcttctt atgatcagct cggtgtgtgt ctcctcctac cgcgggcgca agtcggggaa caagcctccg tccaaaacat gtctgaagga ggag atg gcc aag ggc Met Ala Lys Gly 1</pre>	116
<pre> <211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908) <223> K+Hnov4 <400> 3 ggagccccgc agcgcttctt atgatcagct cggtgtgtgt ctcctcctac cgcgggcgca agtcggggaa caagcctccg tccaaaacat gtctgaagga ggag atg gcc aag ggc Met Ala Lys Gly 1 gag gcg tcg gag aag atc atc atc aac gtg ggc ggc acg cga cat gag Glu Ala Ser Glu Lys Ile Ile Ile Asn Val Gly Gly Thr Arg His Glu 5 10 15 20 </pre>	116
<pre> <211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908) <223> K+Hnov4 <400> 3 ggagccccgc agcgcttctt atgatcagct cggtgtgtgt ctcctcctac cgcgggcgca agtcggggaa caagcctccg tccaaaacat gtctgaagga ggag atg gcc aag ggc Met Ala Lys Gly 1 gag gcg tcg gag aag atc atc atc aac gtg ggc ggc acg cga cat gag Glu Ala Ser Glu Lys Ile Ile Ile Asn Val Gly Gly Thr Arg His Glu 5 10 15 20 acc tac cgc agc acc ctg cgc acc cta ccg gga acc cgc ctc gcc tgg Thr Tyr Arg Ser Thr Leu Arg Thr Leu Pro Gly Thr Arg Leu Ala Trp </pre>	116
<pre> <211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908) <223> K+Hnov4 <400> 3 ggagccccgc agcgcttctt atgatcagct cggtgtgtgt ctcctcctac cgcgggcgca agtcggggaa caagcctccg tccaaaacat gtctgaagga ggag atg gcc aag ggc Met Ala Lys Gly 1 gag gcg tcg gag aag atc atc atc aac gtg ggc ggc acg cga cat gag Glu Ala Ser Glu Lys Ile Ile Ile Asn Val Gly Gly Thr Arg His Glu 5 10 15 20</pre>	116 164 212
<pre></pre>	116
<pre> <211> 1927 <212> DNA <213> H. sapiens <220> <221> CDS <222> (105)(1908) <223> K+Hnov4 <400> 3 ggagccccgc agcgcttctt atgatcagct cggtgtgtgt ctcctcctac cgcgggcgca agtcggggaa caagcctccg tccaaaacat gtctgaagga ggag atg gcc aag ggc Met Ala Lys Gly 1 gag gcg tcg gag aag atc atc atc aac gtg ggc ggc acg cga cat gag Glu Ala Ser Glu Lys Ile Ile Ile Asn Val Gly Gly Thr Arg His Glu 5 10 15 20 acc tac cgc agc acc ctg cgc acc cta ccg gga acc cgc ctc gcc tgg Thr Tyr Arg Ser Thr Leu Arg Thr Leu Pro Gly Thr Arg Leu Ala Trp 35</pre>	116 164 212

gtg ggt agc agc ggc agc ggc ggc ggg ggc tgc gag ttc ttc ttc Val Gly Ser Ser Gly Ser Ser Gly Gly Gly Cys Glu Phe Phe 55 60 65	308
gac agg cac ccg ggc gtc ttc gcc tac gtg ctc aac tac tac cgc acc Asp Arg His Pro Gly Val Phe Ala Tyr Val Leu Asn Tyr Tyr Arg Thr 70 75 80	356
ggc aag ctg cac tgc ccc gca gac gtg tgc ggg ccg ctc ttc gag gag Gly Lys Leu His Cys Pro Ala Asp Val Cys Gly Pro Leu Phe Glu Glu 85 90 95 100	404
gag ctg gcc ttc tgg ggc atc gac gag acc gac gtg gag ccc tgc tgc Glu Leu Ala Phe Trp Gly Ile Asp Glu Thr Asp Val Glu Pro Cys 105 110 115	452
tgg atg acc tac cgg cag cac cgc gac gcc gag gag gcg ctg gac atc Trp Met Thr Tyr Arg Gln His Arg Asp Ala Glu Glu Ala Leu Asp Ile 120 125 130	500
ttc gag acc ccc gac ctc att ggc ggc gac ccc ggc gac gac gag gac Phe Glu Thr Pro Asp Leu Ile Gly Gly Asp Pro Gly Asp Asp Glu Asp 135 140 145	548
ctg gcg gcc aag agg ctg ggc atc gag gac gcg gcg ggg ctc ggg ggc Leu Ala Ala Lys Arg Leu Gly Ile Glu Asp Ala Ala Gly Leu Gly Gly 150 155 160	596
ccc gac ggc aaa tct ggc cgc tgg agg agg ctg cag ccc cgc atg tgg Pro Asp Gly Lys Ser Gly Arg Trp Arg Arg Leu Gln Pro Arg Met Trp 165 170 180	644
gcc ctc ttc gaa gac ccc tac tcg tcc aga gcc gcc agg ttt att gct Ala Leu Phe Glu Asp Pro Tyr Ser Ser Arg Ala Ala Arg Phe Ile Ala 185 190 195	692
ttt gct tct tta ttc ttc atc ctg gtt tca att aca act ttt tgc ctg Phe Ala Ser Leu Phe Phe Ile Leu Val Ser Ile Thr Thr Phe Cys Leu 200 205 210	740
gaa aca cat gaa gct ttc aat att gtt aaa aac aag aca gaa cca gtc Glu Thr His Glu Ala Phe Asn Ile Val Lys Asn Lys Thr Glu Pro Val 215 220 225	788
atc aat ggc aca agt gtt gtt cta cag tat gaa att gaa acg gat cct Ile Asn Gly Thr Ser Val Val Leu Gln Tyr Glu Ile Glu Thr Asp Pro 230 235 240	836
gcc ttg acg tat gta gaa gga gtg tgt gtg gtg tgg ttt act ttt gaa Ala Leu Thr Tyr Val Glu Gly Val Cys Val Val Trp Phe Thr Phe Glu 245 250 255 260	884
ttt tta gtc cgt att gtt ttt tca ccc aat aaa ctt gaa ttc atc aaa Phe Leu Val Arg Ile Val Phe Ser Pro Asn Lys Leu Glu Phe Ile Lys 265 270 275	932
aat ctc ttg aat atc att gac ttt gtg gcc atc cta cct ttc tac tta Asn Leu Leu Asn Ile Ile Asp Phe Val Ala Ile Leu Pro Phe Tyr Leu 280 285 290	980
gag gtg gga ctc agt ggg ctg tca tcc aaa gct gct aaa gat gtg ctt	1028

Glu Val Gly 1	Leu Ser Gl	y Leu Ser 300	Ser Lys Al	a Ala Lys Asp 305	Val Leu
ggc ttc ctc Gly Phe Leu 310	agg gtg gt Arg Val Va	a agg ttt al Arg Phe 315	gtg agg at Val Arg I	cc ctg aga att le Leu Arg Ile 320	ttc aag 1076 Phe Lys
ctc acc cgc Leu Thr Arg 325	His Phe V	ta ggt ctg al Gly Leu 30	Arg val L	tt gga cat act eu Gly His Thr 35	ctt cga 1124 Leu Arg 340
gct agt act Ala Ser Thr	aat gaa t Asn Glu P 345	tt ttg ctg he Leu Leu	ctg ata a Leu Ile I 350	tt ttc ctg gct le Phe Leu Ala	cta gga 1172 Leu Gly 355
gtt ttg ata Val Leu Ile	ttt gct a Phe Ala T 360	cc atg atc hr Met Ile	tac tat g Tyr Tyr A 365	cc gag aga gtg la Glu Arg Val 370	gga gct 1220 Gly Ala
caa cct aac Gln Pro Asn 375	gac cct t Asp Pro S	ca gct agt er Ala Ser 380	GIU HIS I	ca cag ttc aaa hr Gln Phe Lys 385	aac att 1268 Asn Ile
ccc att ggg Pro Ile Gly 390	ttc tgg t	gg gct gta Trp Ala Val 395	gtg acc a Val Thr N	atg act acc ctg Met Thr Thr Leu 400	ggt tat 1316 Gly Tyr
ggg gat atg Gly Asp Met 405	Tyr Pro (caa aca tgg Gln Thr Trp 410	b ser gra t	atg ctg gtg gga Met Leu Val Gly 415	gcc ctg 1364 Ala Leu 420
tgt gct ctg Cys Ala Leu	gct gga g Ala Gly 425	gtg ctg aca Val Leu Th	a ata gcc a r Ile Ala 1 430	atg cca gtg cc Met Pro Val Pro	gtc att 1412 Val Ile 435
gtc aat aat Val Asn Asr	ttt gga Phe Gly 440	atg tac ta Met Tyr Ty	c tcc ttg r Ser Leu 445	gca atg gca aa Ala Met Ala Ly 45	3 0122 -10
ctt cca agg Leu Pro Arg 459	Lys Arg	aag aag ca Lys Lys Hi 46	s lie Pio	cct gct cct ca Pro Ala Pro Gl 465	g gca agc 1508 n Ala Ser
tca cct act Ser Pro Th	t ttt tgc r Phe Cys	aag aca ga Lys Thr Gl 475	a tta aat lu Leu Asn	atg gcc tgc aa Met Ala Cys As 480	t agt aca 1556 n Ser Thr
cag agt ga Gln Ser As 485	c aca tgt p Thr Cys	ctg ggc aa Leu Gly Ly 490	aa gac aat ys Asp Asn	cga ctt ctg ga Arg Leu Leu Gl 495	a cat aac 1604 u His Asn 500
	g tta tca l Leu Ser 505	ggt gac ga Gly Asp As	ac agt aca sp Ser Thr 510	gga agt gag co Gly Ser Glu P	eg cca cta 1652 co Pro Leu 515
tca ccc cc Ser Pro Pr	a gaa agg o Glu Arg 520	ctc ccc a	tc aga cgc le Arg Arg 525	tct agt acc a Ser Ser Thr A	ga gac aaa 1700 rg Asp Lys 30
aac aga ag Asn Arg Ai		aca tgt t Thr Cys P	tc cta ctg he Leu Leu	acg aca ggt g Thr Thr Gly A	at tac acg 1748 sp Tyr Thr

545 540 535 tgt gct tct gat gga ggg atc agg aaa gga tat gaa aaa tcc cga agc 1796 Cys Ala Ser Asp Gly Gly Ile Arg Lys Gly Tyr Glu Lys Ser Arg Ser 555 tta aac aac ata gcg ggc ttg gca ggc aat gct ctg agg ctc tct cca 1844 Leu Asn Asn Ile Ala Gly Leu Ala Gly Asn Ala Leu Arg Leu Ser Pro 570 gta aca tea eec tac aac tet eet tgt eet etg agg ege tet ega tet 1892 Val Thr Ser Pro Tyr Asn Ser Pro Cys Pro Leu Arg Arg Ser Arg Ser 590 585 1927 ccc atc cca tct atc t tgtaaaccaa accctcgtg Pro Ile Pro Ser Ile <210> 4 <211> 601 <212> PRT <213> H. sapiens Met Ala Lys Gly Glu Ala Ser Glu Lys Ile Ile Asn Val Gly Gly <400> 4 10 5 Thr Arg His Glu Thr Tyr Arg Ser Thr Leu Arg Thr Leu Pro Gly Thr 25 20 Arg Leu Ala Trp Leu Ala Asp Pro Asp Gly Gly Gly Arg Pro Glu Thr 40 Asp Gly Gly Gly Val Gly Ser Ser Gly Ser Ser Gly Gly Gly Cys 55 Glu Phe Phe Phe Asp Arg His Pro Gly Val Phe Ala Tyr Val Leu Asn 70 Tyr Tyr Arg Thr Gly Lys Leu His Cys Pro Ala Asp Val Cys Gly Pro 90 85 Leu Phe Glu Glu Glu Leu Ala Phe Trp Gly Ile Asp Glu Thr Asp Val 105 Glu Pro Cys Cys Trp Met Thr Tyr Arg Gln His Arg Asp Ala Glu Glu 115 Ala Leu Asp Ile Phe Glu Thr Pro Asp Leu Ile Gly Gly Asp Pro Gly 140 135 Asp Asp Glu Asp Leu Ala Ala Lys Arg Leu Gly Ile Glu Asp Ala Ala 155 150 Gly Leu Gly Gly Pro Asp Gly Lys Ser Gly Arg Trp Arg Arg Leu Gln 170 Pro Arg Met Trp Ala Leu Phe Glu Asp Pro Tyr Ser Ser Arg Ala Ala 165 185 Arg Phe Ile Ala Phe Ala Ser Leu Phe Phe Ile Leu Val Ser Ile Thr 195 200 205 Thr Phe Cys Leu Glu Thr His Glu Ala Phe Asn Ile Val Lys Asn Lys 220 215 Thr Glu Pro Val Ile Asn Gly Thr Ser Val Val Leu Gln Tyr Glu Ile

250

Glu Thr Asp Pro Ala Leu Thr Tyr Val Glu Gly Val Cys Val Val Trp

Phe Thr Phe Glu Phe Leu Val Arg Ile Val Phe Ser Pro Asn Lys Leu

265 Glu Phe Ile Lys Asn Leu Leu Asn Ile Ile Asp Phe Val Ala Ile Leu 280

230

245

```
Pro Phe Tyr Leu Glu Val Gly Leu Ser Gly Leu Ser Ser Lys Ala Ala
                    295
Lys Asp Val Leu Gly Phe Leu Arg Val Val Arg Phe Val Arg Ile Leu
                                    315
                 310
Arg Ile Phe Lys Leu Thr Arg His Phe Val Gly Leu Arg Val Leu Gly
                       330
His Thr Leu Arg Ala Ser Thr Asn Glu Phe Leu Leu Ile Ile Phe
         340 345
Leu Ala Leu Gly Val Leu Ile Phe Ala Thr Met Ile Tyr Tyr Ala Glu
                         360
Arg Val Gly Ala Gln Pro Asn Asp Pro Ser Ala Ser Glu His Thr Gln
                                       380
                    375
Phe Lys Asn Ile Pro Ile Gly Phe Trp Trp Ala Val Val Thr Met Thr
                             395
                  390
Thr Leu Gly Tyr Gly Asp Met Tyr Pro Gln Thr Trp Ser Gly Met Leu 405 410 415
Val Gly Ala Leu Cys Ala Leu Ala Gly Val Leu Thr Ile Ala Met Pro
                     425
 Val Pro Val Ile Val Asn Asn Phe Gly Met Tyr Tyr Ser Leu Ala Met
         420
                   440
 Ala Lys Gln Lys Leu Pro Arg Lys Arg Lys Lys His Ile Pro Pro Ala
      435
                     455
 Pro Gln Ala Ser Ser Pro Thr Phe Cys Lys Thr Glu Leu Asn Met Ala
                            . 475
         470
 Cys Asn Ser Thr Gln Ser Asp Thr Cys Leu Gly Lys Asp Asn Arg Leu
485 490 495
 Leu Glu His Asn Arg Ser Val Leu Ser Gly Asp Asp Ser Thr Gly Ser
                             505
 Glu Pro Pro Leu Ser Pro Pro Glu Arg Leu Pro Ile Arg Arg Ser Ser
           500
                                           525
        515
                         520
 Thr Arg Asp Lys Asn Arg Arg Gly Glu Thr Cys Phe Leu Leu Thr Thr
                                         540
             535
 Gly Asp Tyr Thr Cys Ala Ser Asp Gly Gly Ile Arg Lys Gly Tyr Glu
                                   555
            550
 Lys Ser Arg Ser Leu Asn Asn Ile Ala Gly Leu Ala Gly Asn Ala Leu
                               570
             565
 Arg Leu Ser Pro Val Thr Ser Pro Tyr Asn Ser Pro Cys Pro Leu Arg
                            585
          580
  Arg Ser Arg Ser Pro Ile Pro Ser Ile
         595
       <210> 5
        <211> 2293
        <212> DNA
        <213> H. sapiens
        <220>
        <221> CDS
        <222> (330) ... (1800)
        <223> K+Hnov6
   gggaagagcg aacccagggc cettgetete gtgcageget gegeeetggg tggggaegge
   gtgaggettg cagegeaggt gagagtgatt ttecagtgat tgetttggee tgtacaacca
                                                                   120
   gagaacagga ttettecett etttttggcc accaaatgce tatgtgcacc acacatteca
                                                                   180
   gtgtgctgag aagggcagag cttcttggat gatgatggac gtcccaccgg gcaggatgaa
                                                                   240
   ggcagagcgt gtggcatctc cacctcaagg gtgcagcctg atcttcctct tctcccttgc
                                                                   300
   cagccagcac tetgeettet gtatecace atg gtg ttt ggt gag ttt tte cat
                                                                   353
                                Met Val Phe Gly Glu Phe Phe His
                                 1
```

cgc cct gga caa gac gag gaa ctt gtc aac ctg aat gtg ggg ggc ttt Arg Pro Gly Gln Asp Glu Glu Leu Val Asn Leu Asn Val Gly Gly Phe 10 15 20	401
aag cag tot gtt gac caa agc acc oto otg ogg ttt cot cac acc aga Lys Gln Ser Val Asp Gln Ser Thr Leu Leu Arg Phe Pro His Thr Arg 25 30 35 40	449
ctg ggg aag ctg ctt act tgc cat tct gaa gag gcc att ctg gag ctg Leu Gly Lys Leu Leu Thr Cys His Ser Glu Glu Ala Ile Leu Glu Leu 45 50 55	497
tgt gat gat tac agt gtg gcc gat aag gaa tac tac ttt gat cgg aat Cys Asp Asp Tyr Ser Val Ala Asp Lys Glu Tyr Tyr Phe Asp Arg Asn 60 65 70	545
ccc tcc ttg ttc aga tat gtt ttg aat ttt tat tac acg ggg aag ctg Pro Ser Leu Phe Arg Tyr Val Leu Asn Phe Tyr Tyr Thr Gly Lys Leu 75 80 85	593
cat gtc atg gag gag ctg tgc gta ttc tca ttc tgc cag gag atc gag His Val Met Glu Glu Leu Cys Val Phe Ser Phe Cys Gln Glu Ile Glu 90 95 100	641
tac tgg ggc atc aac gag ctc ttc att gat tct tgc tgc agc aat cgc Tyr Trp Gly Ile Asn Glu Leu Phe Ile Asp Ser Cys Cys Ser Asn Arg 105 110 115 120	689
tac cag gaa cgc aag gag gaa aac cac gag aag gac tgg gac cag aaa Tyr Gln Glu Arg Lys Glu Glu Asn His Glu Lys Asp Trp Asp Gln Lys 125 130 135	737
agc cat gat gtg agt acc gac tcc tcg ttt gaa gag tcg tct ctg ttt Ser His Asp Val Ser Thr Asp Ser Ser Phe Glu Glu Ser Ser Leu Phe 140 145 150	785
gag aaa gag ctg gag aag ttt gac aca ctg cga ttt ggt cag ctc cgg Glu Lys Glu Leu Glu Lys Phe Asp Thr Leu Arg Phe Gly Gln Leu Arg 155 160 165	833
aag aaa atc tgg att aga atg gag aat cca gcg tac tgc ctg tcc gct Lys Lys Ile Trp Ile Arg Met Glu Asn Pro Ala Tyr Cys Leu Ser Ala 170 175 180	881
aag ctt atc gct atc tcc tcc ttg agc gtg gtg ctg gcc tcc atc gtg Lys Leu Ile Ala Ile Ser Ser Leu Ser Val Val Leu Ala Ser Ile Val 190 195 200	929
gcc atg tgc gtt cac agc atg tcg gag ttc cag aat gag gat gga gaa Ala Met Cys Val His Ser Met Ser Glu Phe Gln Asn Glu Asp Gly Glu 205 210	977
	1025
gtg gat gat ccg gtg ctg gaa gga gtg gag atc gcg tgc att gcc tgg Val Asp Asp Pro Val Leu Glu Gly Val Glu Ile Ala Cys Ile Ala Trp 220 225 230	1025
Val Asp Asp Pro Val Leu Giu Giy Val Gia 110	1073

Lys Phe Trp Lys Asn Pro Leu Asn Ile Ile Asp Phe Val Ser Ile Ile 250 255 260	
ccc ttc tat gcc acg ttg gct gta gac acc aag gag gaa gag agt gag Pro Phe Tyr Ala Thr Leu Ala Val Asp Thr Lys Glu Glu Glu Ser Glu 265 270 280	1169
gat att gag aac atg ggc aag gtg gtc cag atc cta cgg ctt atg agg Asp Ile Glu Asn Met Gly Lys Val Val Gln Ile Leu Arg Leu Met Arg 285 290 295	1217
att ttc cga att cta aag ctt gcc cgg cac tcg gta gga ctt cgg tct Ile Phe Arg Ile Leu Lys Leu Ala Arg His Ser Val Gly Leu Arg Ser 300 305 310	1265
cta ggt gcc aca ctg aga cac agc tac cat gaa gtt ggg ctt ctg ctt Leu Gly Ala Thr Leu Arg His Ser Tyr His Glu Val Gly Leu Leu Leu 315 320 325	1313
ctc ttc ctc tct gtg ggc att tcc att ttc tct gtg ctt atc tac tcc Leu Phe Leu Ser Val Gly Ile Ser Ile Phe Ser Val Leu Ile Tyr Ser 330 335 340	1361
gtg gag aaa gat gac cac aca tcc agc ctc acc agc atc ccc atc tgc Val Glu Lys Asp Asp His Thr Ser Ser Leu Thr Ser Ile Pro Ile Cys 345 350 355 360	1409
tgg tgg tgg gcc acc atc agc atg aca act gtg ggc tat gga gac acc Trp Trp Trp Ala Thr Ile Ser Met Thr Thr Val Gly Tyr Gly Asp Thr 365 370 375	1457
cac ccg gtc acc ttg gcg gga aag ctc atc gcc agc aca tgc atc atc His Pro Val Thr Leu Ala Gly Lys Leu Ile Ala Ser Thr Cys Ile Ile 380 385 390	1505
tgt ggc atc ttg gtg gtg gcc ctt ccc atc acc atc atc ttc aac aag Cys Gly Ile Leu Val Val Ala Leu Pro Ile Thr Ile Ile Phe Asn Lys 395 400 405	1553
ttt tcc aag tac tac cag aag caa aag gac att gat gtg gac cag tgc Phe Ser Lys Tyr Tyr Gln Lys Gln Lys Asp Ile Asp Val Asp Gln Cys 410 415 420	1601
agt gag gat gca cca gag aag tgt cat gag cta cct tac ttt aac att Ser Glu Asp Ala Pro Glu Lys Cys His Glu Leu Pro Tyr Phe Asn Ile 425 430 435 440	1649
agg gat ata tat gca cag cgg atg cac gcc ttc att acc agt ctc tct Arg Asp Ile Tyr Ala Gln Arg Met His Ala Phe Ile Thr Ser Leu Ser 445 455	1697
tct gta ggc att gtg gtg agc gat cct gac tcc aca gat gct tca agc Ser Val Gly Ile Val Val Ser Asp Pro Asp Ser Thr Asp Ala Ser Ser 460 465 470	1745
att gaa gac aat gag gac att tgt aac acc acc tcc ttg gag aat tgc Ile Glu Asp Asn Glu Asp Ile Cys Asn Thr Thr Ser Leu Glu Asn Cys 475 480 485	1793
aca gca a aatgagcggg ggtgtttgtg cetgtttctc ttatcctttc ccaacattag Thr Ala	1850

2293

490

gttaacacag ctttataaac ctcagtgggt tcgttaaaat catttaattc tcagggtgta cctttcagcc atagttggac attcattgct gaattctgaa atgatagaat tgtctttatt 1970 tttctctgtg aggtcaatta aatgccttgt tctgaaattt atttttaca agagagagtt 2030 gtgatagagt ttggaatata agataaatgg tattgggtgg ggtttgtggc tacagcttat gcatcattct gtgtttgtca tttactcaca ttgagctaac tttaaattac tgacaagtag 2150 aatcaaaggt gcagctgact gagacgacat gcatgtaaga tccacaaaat gagacaatgc 2210 atgtaaatcc atgctcatgt tctaaacatg gaaactagga gcctaataaa cttcctaatt 2270 .cagaaaaaaa aaaaaaaaaa aaa <210> 6 <211> 490 <212> PRT <213> H. sapiens Met Val Phe Gly Glu Phe Phe His Arg Pro Gly Gln Asp Glu Glu Leu <400> 6 1 5 Val Asn Leu Asn Val Gly Gly Phe Lys Gln Ser Val Asp Gln Ser Thr 25 Leu Leu Arg Phe Pro His Thr Arg Leu Gly Lys Leu Leu Thr Cys His 40 Ser Glu Glu Ala Ile Leu Glu Leu Cys Asp Asp Tyr Ser Val Ala Asp 35 55 Lys Glu Tyr Tyr Phe Asp Arg Asn Pro Ser Leu Phe Arg Tyr Val Leu 70 Asn Phe Tyr Tyr Thr Gly Lys Leu His Val Met Glu Glu Leu Cys Val 90 85 Phe Ser Phe Cys Gln Glu Ile Glu Tyr Trp Gly Ile Asn Glu Leu Phe 105 Ile Asp Ser Cys Cys Ser Asn Arg Tyr Gln Glu Arg Lys Glu Glu Asn 115 120 125 100 His Glu Lys Asp Trp Asp Gln Lys Ser His Asp Val Ser Thr Asp Ser 140 135 Ser Phe Glu Glu Ser Ser Leu Phe Glu Lys Glu Leu Glu Lys Phe Asp 155 Thr Leu Arg Phe Gly Gln Leu Arg Lys Lys Ile Trp Ile Arg Met Glu 150 170 165 Asn Pro Ala Tyr Cys Leu Ser Ala Lys Leu Ile Ala Ile Ser Ser Leu 185 Ser Val Val Leu Ala Ser Ile Val Ala Met Cys Val His Ser Met Ser 200 Glu Phe Gln Asn Glu Asp Gly Glu Val Asp Asp Pro Val Leu Glu Gly 220 215 Val Glu Ile Ala Cys Ile Ala Trp Phe Thr Gly Glu Leu Ala Val Arg 230 Leu Ala Ala Pro Cys Gln Lys Lys Phe Trp Lys Asn Pro Leu Asn 250 245 Ile Ile Asp Phe Val Ser Ile Ile Pro Phe Tyr Ala Thr Leu Ala Val 265 Asp Thr Lys Glu Glu Glu Ser Glu Asp Ile Glu Asn Met Gly Lys Val 285 280 Val Gln Ile Leu Arg Leu Met Arg Ile Phe Arg Ile Leu Lys Leu Ala 295 Arg His Ser Val Gly Leu Arg Ser Leu Gly Ala Thr Leu Arg His Ser

330

Tyr His Glu Val Gly Leu Leu Leu Phe Leu Ser Val Gly Ile Ser

Ile Phe Ser Val Leu Ile Tyr Ser Val Glu Lys Asp Asp His Thr Ser

345

310

325

Ser Leu Thr Ser Ile Pro Ile Cys Trp Trp Trp Ala Thr Ile Ser Met	
Thr Thr Val Gly Tyr Gly Asp Thr His Pro Val Thr Leu Ala Gly Lys	
270 375 Leu Ile Ala Ser Thr Cys Ile Ile Cys Gly Ile Leu Val Val Ala Leu 395 400	
395 390 395 The Clarke Gla	
385 390 Pro Ile Thr Ile Ile Phe Asn Lys Phe Ser Lys Tyr Tyr Gln Lys Gln 415 405 410	
Lys Asp Ile Asp Val Asp Gln Cys Ser Glu Asp Ala Pro Glu Lys Cys 430 425 430	
420 425 His Glu Leu Pro Tyr Phe Asn Ile Arg Asp Ile Tyr Ala Gln Arg Met 445 440 445	
His Ala Phe Ile Thr Ser Leu Ser Ser Val Gly Ile Val Val Ser Asp 455 460	
450 455 Pro Asp Ser Thr Asp Ala Ser Ser Ile Glu Asp Asn Glu Asp Ile Cys 475 480	
4/0	
Asn Thr Thr Ser Leu Glu Asn Cys Thr Ala 485 490	
<210> 7 <211> 3080	
<211> DNA	
<213> H. sapiens	
<220>	
<221> CDS <222> (480)(1977)	
<223> K+Hnov9	
<400> 7 gtctctcctc ttcctcctcc tccgccccac atctccctc ttcctccctt ccccaacccc gtctctcctc ttcctcctcc tccgccccac trtacacctc ctggcctggg aatcgcaatt	60
gteteteete tteeteete teegeeeta atteeteete tegggetggg aategeaatt teeaeceae aagtagegag teatteaate tgtacaecete etgggetggg aategeaatt	120 180
tecacecace aagtagegag teatteaate tytacacete eegsperson geagtgtgca gegaagttgg gaggeggggt gacaacettt gggaagtgtee agggegaceg geagtgtgca gegaagttgg gagagttge eestteetet	
	2/10
gcgaagttgg gaggcgggg gacctcacct gatcctctct cttagcgcga cccttcctct	240 300
cagggactgt gtegggettg gaeettatte gestaattes greactee geteetag	300
cagggactgt gtcgggcttg gaccttattt gaccgcgcttcc gcgcactccc ggctccctag gctccctgtc tcctctttct gccacttgtg cgctgcttcc gcgcactccc ggtaacccct	
cagggactgt gtcgggcttg gaccttactt gaccgcgttcc gcgcactccc ggctccctag gctccctgtc tcctctttct gccacttgtg cgctgcttcc gcgcactccc ggctccctt cggcaggagg aggaaggcgc acagcgggtg gagagggtgc gccaaggaga ggtaacccct	300 360
cagggactgt gtegggettg gacettaett gacegtgettee gegeaeteee ggeteeetag geteeetgte teetetttet gecaettgtg egetgettee gegeaaggag ggtaaceeet eggeaggagg aggaaggege acagegggtg gagaagggtge gecaaeggaga ggtaaceeet eggggageee ggggaateee ggeegeeaee aggggeegtg ecaeegeeet egegggacea teggggageee ggggaateee ggeegeeaee aggggeegtg ecaeegeeet gggttagag	300 360 420
cagggactgt gtegggettg gacettaett gaceggactec ggetecetag getecetagte teetetttet gecaettgtg egtgettec gegeaettec ggetaecet egggagggagg aggaaggege acagegggtg gagagggtge gecaaggaga ggtaaceet tegggagece ggggaateee ggeegecace aggggeegtg egegeeet egegggacta gggttagag aagetteegg egtgteeea aetttgtgge geeeteagge egeggegact gggttagag	300 360 420 479
cagggactgt gtegggettg gacettaett gatestette gegeactece ggeteeetag geteeetgte teetetttet gecacttgtg egetgettee gegeactece ggetaeeet eggeaggagg aggaaggege acagegggtg gagagggtge gecaaggaga ggtaaceeet tegggagece ggggaateee ggeegecace aggggeegtg ecacegeeet egegggacea aagetteegg egtgteeea aetttgtgge geeeteagge egeggegact gggttagag atg eet tee age gge aga geg etg etg gae teg eeg etg gae age gge atg eet tee age gge Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly Met Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly	300 360 420 479
cagggactgt gtegggettg gacettaett gegegettee gegeactee ggeteectag geteectgte teetetttet gecacttgtg egtgettee gegeactee ggetaecet eggeaggagg aggaaggege acagegggtg gagagggtge gecaaggaga ggtaaceeet tegggagece ggggaateee ggeegecace aggggeegtg ecacegeeet eggggacea aagetteegg egtgteecea actttgtgge geceteagge egeggegact gggttagag atg eet tee age gge aga geg etg etg gae teg eeg etg gae age gge Met Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly	300 360 420 479 527
cagggactgt gtegggettg gacettaett gtegggettee gegeactee ggeteectag geteectgte teetettet gecaettgtg egetgettee gegeactee ggetaaceet eggeaggagg aggaaggeg acagegggtg gagagggtge gecaaggaga ggtaaceet tegggagee ggggaateee ggeegecace aggggeegtg eeaeegeeet egegggacea aagetteegg egtgteecea aetttgtgge geeeteagge egeggegact gggttagag atg cet tee age gge aga geg etg etg gae teg eeg etg gae age gge Met Pro Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly 15 15 16 15 15	300 360 420 479
cagggactgt gtegggettg gacettaett geagetgettee gegeactee ggeteectag geteectgte teetettet gecaettgtg egetgettee gegeactee ggetaaceet eggeaggagg aggaaggege acaaggggtg gagagggtge gecaaggaga ggtaaceet tegggageee ggggaatee ggeegeace aggggeegtg ecaeegeeet egegggacea aagetteegg egtgteecea actttgtgge geecteagge egeggegaet gggttagag atg eet tee age gge aga geg etg etg gae teg eeg etg gae age gge Met Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly 1 1 5 15 tee etg ace tee etg gae tet agt gte tte tge age gag ggt gaa ggg Ser Leu Thr Ser Leu Asp Ser Ser Val Phe Cys Ser Glu Gly Glu Gly	300 360 420 479 527
cagggactgt gtegggettg gacettaett geacettaett gegeactee ggeteettag geteettet tetettett geacettgtg egetgettee gegeactee ggetaaceet eggeaggagg aggaaggegg acaaggaggg gagagggtge gecaaggaga ggtaaceet tegggagee ggggaatee ggegeaceae aggggeegtg ecaeegeeet egegggacea aagetteegg egtgteecaa actttgtgge geeetcagge egeggegaet gggttagag atg eet tee age gge aga geg etg etg gae teg eeg etg gae age gge Met Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly 1 1 5 15 tee etg ace tee etg gae tet agt gte tte tge age gag ggt gaa ggg Ser Leu Thr Ser Leu Asp Ser Ser Val Phe Cys Ser Glu Gly Glu Gly 20 25 30	300 360 420 479 527
cagggactgt gtegggettg gacettaett geagetgettee gegeactee ggeteectag geteectgte teetettet gecaettgtg egetgettee gegeactee ggetaaceet eggeaggag aggaaggege acaaggggtg gagagggtge gecaaggaga ggtaaceet tegggagee ggggaatee ggeegeace aggggeegtg ecaeegeeet egegggacea aagetteegg egtgteecea actttgtgge geeetcagge egeggegaet gggttagag atg eet tee age gge aga geg etg etg gae teg eeg etg gae age gge atg eeg etg gae age gge 10 leu Asp Ser Gly Met Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly 10 lece etg ace etg gae tee etg age gag ggt gaa ggg ser Leu Thr Ser Leu Asp Ser Ser Val Phe Cys Ser Glu Gly Glu Gly 20 lece etg age age gge age	300 360 420 479 527
cagggactgt gtegggettg gacettaett geagetgettee gegeactee ggeteectag geteectgte teetettet gecaettgtg egetgettee gegeactee ggetaaceet eggeaggag aggaaggege acaaggggtg gagagggtge gecaaggaga ggtaaceet tegggagee ggggaatee ggeegeace aggggeegtg ecaeegeeet egegggacea aagetteegg egtgteecea actttgtgge geeetcagge egeggegaet gggttagag atg eet tee age gge aga geg etg etg gae teg eeg etg gae age gge atg eeg etg gae age gge 10 leu Asp Ser Gly Met Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly 10 lece etg ace etg gae tee etg age gag ggt gaa ggg ser Leu Thr Ser Leu Asp Ser Ser Val Phe Cys Ser Glu Gly Glu Gly 20 lece etg age age gge age	300 360 420 479 527
cagggactgt gtegggettg gacettaett geagetgettee gegeactee ggeteectag geteectgte teetettet geaacttgtg egetgettee gegeaggag aggaaggega acaaggggtg gagagggtge geaaggaga ggtaaceet tegggagee ggggaatee ggcegeace aggggegtg ecaeggeet egegggacea aagetteegg egtgteecea actttgtgge geeetcagge egeggegaet gggttagag atg eet tee age gge aga geg etg etg gae teg eeg etg gae age gge Met Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly 1 1 5 15 tee etg ace tee etg gae tet agt gte tte tge age gag ggt gaa ggg ser Leu Thr Ser Leu Asp Ser Ser Val Phe Cys Ser Glu Gly Glu Gly gag eee ttg geg ete ggg gae tge tte aeg gte aac gtg gge gge age Glu Pro Leu Ala Leu Gly Asp Cys Phe Thr Val Asn Val Gly Gly Ser	300 360 420 479 527
cagggactgt gtegggettg gacettaett geagetgettee gegeactee ggeteectag geteectge teetettet geaacttgtg egetgettee gegeaggag aggaaggega acaaggggtg gagaagggtge geaaggaga ggtaaceet tegggagee ggggaatee ggcegeace aggggegtg ecaeggeet egegggacea aagetteegg egtgteecea actttgtgge geeetcagge egeggegaet gggttagag atg eet tee age gge aga geg etg etg gae teg eeg etg gae age gge Met Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly 1	300 360 420 479 527 575
cagggactgt gtegggettg gacettaett geaggtegettee gegeactee ggeteectag geteectgte teetettet geaacttgtg egetgettee gegeaggag aggaaggege acaaggggtg gagagggtge geaaggaga ggtaaceet tegggagee ggggaatee ggeegeace aggggegtg ecaaeggeet egegggacea aagetteegg egtgteecea actttgtgge geeteagge egeggegaet gggttagag atg eet tee age gge aga geg etg etg gae teg eeg etg gae age gge atg eeg etg gae age gge het Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly 10 15 tee etg ace tee etg gae tet agt gte tee tge age gag ggt gaa ggg Ser Leu Thr Ser Leu Asp Ser Ser Val Phe Cys Ser Glu Gly Glu Gly 20 25 30 gag eee ttg geg ete ggg gae tge tte aeg gte aae gtg gge gge age Glu Pro Leu Ala Leu Gly Asp Cys Phe Thr Val Asn Val Gly Gly Ser 40 45	300 360 420 479 527
cagggactgt gtegggettg gacettaett gcacettaett gegeteetaet geteetaet geteetaet geteetaet gegeteetaet gegeteetaetaet gegeteetaetaet gegeteetaetaet gegeteetaetaet gegeteetaetaetaetaetaetaetaetaetaetaetaetae	300 360 420 479 527 575
cagggactgt gtegggettg gacettaett gcacettaett agggactaet gcacetaettaett agggactaettaett gcacettaett gcacettaett agggactaettaett gcacettaett gcacettaett agggactaettaett gcacettaett gcacettaett agggactaettaett gcacettaett gggactaettaett gggactaettaett gcacettaett gggactaettaett gggactaettaett gggactaettaett gggactaettaettaettaettaettaettaettaettaetta	300 360 420 479 527 575
cagggactgt gtegggettg gacettaett gcacettaett gcacettaettaettaettaettaettaettaettaettae	300 360 420 479 527 575
cagggactgt gtegggettg gacettaett gcacettaett gcacettaettaettaettaettaettaettaettaettae	300 360 420 479 527 575 623
cagggactgt gtegggettg gacettaett gcacettgtg geteettete gegeactee ggeteettete gegeacteet geteettete geacettgtg geteettete gegeacteet gegeacteet ggagagggg aggaaggeg aggaaggggg gagagaggtge gecaaggaga ggtaacceet tegggagece ggggaateet ggegeactee aggggeetteegg gagaggggge gecaaggaga ggtaacceet aggegeetteegg ggggaateet ggggtaetee aggggeetteegg gecetcagge egeggegact gggttagag atg cet tee age gge aga geg etg etg gae teg eeg etg gae age gge Met Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly 1	300 360 420 479 527 575 623
cagggactgt gtegggettg gacettaett gcacettaett gcacettaettaett gcacettaettaettaettaettaettaettaettaettae	300 360 420 479 527 575 623
cagggactgt gtegggettg gacettaett gcacettgtg cyctectaet gegeaettee gegeaetteeggeaetgeaetgeaetteeggeaetteeggeaetteeggeaetgeaetgeaetteeggeaetteeggeaetgeae	300 360 420 479 527 575 623
cagggactgt gtegggettg gacettaett gcacettgtg geteettete gegeactee ggeteettete gegeacteet geteettete geacettgtg geteettete gegeacteet gegeacteet ggagagggg aggaaggeg aggaaggggg gagagaggtge gecaaggaga ggtaacceet tegggagece ggggaateet ggegeactee aggggeetteegg gagaggggge gecaaggaga ggtaacceet aggegeetteegg ggggaateet ggggtaetee aggggeetteegg gecetcagge egeggegact gggttagag atg cet tee age gge aga geg etg etg gae teg eeg etg gae age gge Met Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly 1	300 360 420 479 527 575 623

gtg gac aac gag tac ttc ttc gac cgc agc tcg cag gcg ttc cga tat Val Asp Asn Glu Tyr Phe Phe Asp Arg Ser Ser Gln Ala Phe Arg Tyr 100 105 110	815
gtc ctg cac tac tac cgc acc ggc ctg ctg cat gtc atg gag cag ctg Val Leu His Tyr Tyr Arg Thr Gly Arg Leu His Val Met Glu Gln Leu 115 120 125	863
tgc gcg ctc tcc ttc ctg cag gag atc cag tac tgg ggc atc gat gag Cys Ala Leu Ser Phe Leu Gln Glu Ile Gln Tyr Trp Gly Ile Asp Glu 130 135 140	911
ctc agc atc gat tcc tgc tgc agg gac aga tac ttc aga agg aaa gag Leu Ser Ile Asp Ser Cys Cys Arg Asp Arg Tyr Phe Arg Arg Lys Glu 145 150 155 160	959
ctg agt gaa act tta gac ttc aag aag gac aca gaa gac cag gaa agt Leu Ser Glu Thr Leu Asp Phe Lys Lys Asp Thr Glu Asp Gln Glu Ser 165 170 175	1007
caa cat gag agt gaa cag gac ttc tcc caa gga cct tgt ccc act gtt Gln His Glu Ser Glu Gln Asp Phe Ser Gln Gly Pro Cys Pro Thr Val 180 185 190	1055
cgc cag aag ctc tgg aat atc ctg gag aaa cct gga tct tcc aca gct Arg Gln Lys Leu Trp Asn Ile Leu Glu Lys Pro Gly Ser Ser Thr Ala 195 200 205	1103
gcc cgt atc ttt ggc gtc atc tcc att atc ttc gtg gtg gtg tcc atc Ala Arg Ile Phe Gly Val Ile Ser Ile Ile Phe Val Val Val Ser Ile 210 215 220	1151
att aac atg gcc ctg atg tca gct gag tta agc tgg ctg gac ctg cag Ile Asn Met Ala Leu Met Ser Ala Glu Leu Ser Trp Leu Asp Leu Gln 225 230 235 240	1199
ctg ctg gaa atc ctg gag tat gtg tgc att agc tgg ttc acc ggg gag Leu Leu Glu Ile Leu Glu Tyr Val Cys Ile Ser Trp Phe Thr Gly Glu 245 250 255	1247
ttt gtc ctc cgc ttc ctg tgt gtg cgg gac agg tgt cgc ttc cta aga Phe Val Leu Arg Phe Leu Cys Val Arg Asp Arg Cys Arg Phe Leu Arg 260 265 270	1295
aag gtg cca aac atc ata gac ctc ctt gcc atc ttg ccc ttc tac atc Lys Val Pro Asn Ile Ile Asp Leu Leu Ala Ile Leu Pro Phe Tyr Ile 275 280 285	1343
act ctt ctg gta gag agc cta agt ggg agc cag acc acg cag gag ctg Thr Leu Leu Val Glu Ser Leu Ser Gly Ser Gln Thr Thr Gln Glu Leu 290 295 300	1391
gag aac gtg ggg cgc att gtc cag gtg ttg agg ctg ctc agg gct ctg Glu Asn Val Gly Arg Ile Val Gln Val Leu Arg Leu Leu Arg Ala Leu 320 315	1439
cgc atg cta aag ctg ggc aga cat tcc aca gga tta cgc tcc ctt ggg Arg Met Leu Lys Leu Gly Arg His Ser Thr Gly Leu Arg Ser Leu Gly 325 330 335	1487

atg aca atc acc cag tgt tac gaa gaa gtc ggc cta ctg ctc cta ttt 15 Met Thr Ile Thr Gln Cys Tyr Glu Glu Val Gly Leu Leu Leu Phe 340 345 350	535
cta tcc gtg gga atc tct ata ttt tca act gta gaa tac ttt gct gcs Leu Ser Val Gly Ile Ser Ile Phe Ser Thr Val Glu Tyr Phe Ala Glu 355 360 365	5 83
caa agc att cct gac aca acc ttc aca agt gtc cct tgt gca tgg tgg Gln Ser Ile Pro Asp Thr Thr Phe Thr Ser Val Pro Cys Ala Trp Trp 370 375 380	631
	.679
gac acc acc aca ggc aaa atc gtg gcc ttc atg tgt ata tta tcg gga 1 Asp Thr Thr Gly Lys Ile Val Ala Phe Met Cys Ile Leu Ser Gly 405 410 415	L 72 7
att ctt gtc ttg gcc ttg cct att gct att att aac gat cgc ttc tct Ile Leu Val Leu Ala Leu Pro Ile Ala Ile Ile Asn Asp Arg Phe Ser 420 425 430	1775
gct tgc tac ttc acc ttg aaa ctc aag gaa gca gct gtt aga cag cgt Ala Cys Tyr Phe Thr Leu Lys Leu Lys Glu Ala Ala Val Arg Gln Arg 435 440 445	1823
	1871
agt gtt aac ttg aga gat gtc tat gcc cgg agt atc atg gag atg ctg Ser Val Asn Leu Arg Asp Val Tyr Ala Arg Ser Ile Met Glu Met Leu 465 470 475 480	1919
cga ctg aaa ggc aga gaa aga gca agt act agg agc agc ggg gga gat cga ctg aaa ggc aga gaa aga gca agt act agg agc agc ggg gga gat Arg Leu Lys Gly Arg Glu Arg Ala Ser Thr Arg Ser Ser Gly Gly Asp 495 485 490	1967
gat ttc tgg t tttgaattaa ttttcaattt atttacaaaa gctatgtaca Asp Phe Trp	2017
b. sh b a 2	2077
attaactaaa atgataaagc agtgatgtgg atttctgtat tctgatgatg agtctcttca	2137
attaactaaa atgataaagc agtgatgtgg attictgtat todactact agaatatttc gagtactgct catcttaatt aattittgct gatatattgg ttcatctact agaatatttc	2197
gagtactgct catcttaatt aatttttgct gatatattg agtgtccaaa atagccaatt acatcaccta taacaactgc acagtgttct gacacatttg agtgtccaaa tataaaataa	2257
aacacaacca aatacaactg ggccaatata gatttatgt atcactaaca ttagaagttt	2317
tgttattgca atacatacaa aaaayttaaa gattaaagt tagcccagag aaagataagt	2377
tttgcaccac taattttta aaaatggaag stattatcca agtacataaa	2437
aaatatttaa gaacatattg aacaacttty taatataata cttagcttta caagagaaaa	2497
ttactocgtt ctctatcagt taaagctatt gaatataata totagstaa ccactggtca cccatatttg atgggcagag attatatccc tatcttcttt ttcatgtaaa ccactggtca cccatatttg atgggcagag attatatccc tatcttcttt ttcatgtaaa ccactggtca	2557
cccatatttg atgggcagag attatatccc tatettett tetagaattgcatcaaattgct caaatgaact gatctctgta teccattatt actataagag gtggggaatcc caaaactgct caaatgaact gatctctgta teccatatt actataagag gtgggcatctt cattetecca	2617 2677
caaatgaact gatctctgta tcccattatt actataagag gtgsgsact tagattgcag tacatgagtc tacacaaaga cttcaacaat tgcacatctt cattctccca tagattgcag tacatgagtc tacacaaaga catatttctt agtatttcat gaatatcaga	2737
tagattgcag tacatgagte tacacaaaaga etttaataat tagtatttcat gaatatcaga actgagtgta gtatgtggag cataaaaacag catatttctt agtatttcat gaatatcaga	2797
actgagtgta gtatgtggag cataaaacag catatttett agutttaa aataatgaat tggtctttaa atgtctcttt atggatgtat tgttcacatt atggctttaa aataatctta	2857
tggtctttaa atgtctcttt atggatgtat tgttcatatt uggattacta aataatctta atgtaaaagt gaggtagtga acatcctaaa tttctacact ggaattacta aataatctta atgtaaaagt gaggtagtga catcctaaa catcactga tgaacttgaa gatcttttac	2917
atgtaaaagt gaggtagtga acatcctaaa tttttactgga tgaacttgaa gatcttttac tttcataaat gggaaatata tgttaaatga catcactgga tgaacttgaa gatcttttac	2977
tttcataaat gggaaatata tgttaaatg catcattgga tgtatabus s ttgttaacaa aaaaatacta tgggacagctt tctgattgt ggggtaaata gcaaatgttc	3037
aaactttgca ggcattttga Catteateat aacadoudus	
14	

<210> 8
<211> 499
<212> PRT
<213> H. sapiens

<213> H. sapiens <400> 8 Met Pro Ser Ser Gly Arg Ala Leu Leu Asp Ser Pro Leu Asp Ser Gly Ser Leu Thr Ser Leu Asp Ser Ser Val Phe Cys Ser Glu Gly Glu Gly Glu Pro Leu Ala Leu Gly Asp Cys Phe Thr Val Asn Val Gly Gly Ser Arg Phe Val Leu Ser Gln Gln Ala Leu Ser Cys Phe Pro His Thr Arg Leu Gly Lys Leu Ala Val Val Val Ala Ser Tyr Arg Arg Pro Gly Ala 65 70 75 80 Leu Ala Ala Val Pro Ser Pro Leu Glu Leu Cys Asp Asp Ala Asn Pro Val Asp Asn Glu Tyr Phe Phe Asp Arg Ser Ser Gln Ala Phe Arg Tyr Val Leu His Tyr Tyr Arg Thr Gly Arg Leu His Val Met Glu Gln Leu Cys Ala Leu Ser Phe Leu Gln Glu Ile Gln Tyr Trp Gly Ile Asp Glu 130 135 140 Leu Ser Ile Asp Ser Cys Cys Arg Asp Arg Tyr Phe Arg Arg Lys Glu Leu Ser Glu Thr Leu Asp Phe Lys Lys Asp Thr Glu Asp Gln Glu Ser Glm His Glu Ser Glu Gln Asp Phe Ser Gln Gly Pro Cys Pro Thr Val Arg Gln Lys Leu Trp Asn Ile Leu Glu Lys Pro Gly Ser Ser Thr Ala Ala Arg Ile Phe Gly Val Ile Ser Ile Ile Phe Val Val Ser Ile Ile Asn Met Ala Leu Met Ser Ala Glu Leu Ser Trp Leu Asp Leu Gln Leu Leu Glu Ile Leu Glu Tyr Val Cys Ile Ser Trp Phe Thr Gly Glu Phe Val Leu Arg Phe Leu Cys Val Arg Asp Arg Cys Arg Phe Leu Arg Lys Val Pro Asn Ile Ile Asp Leu Leu Ala Ile Leu Pro Phe Tyr Ile 275 280 285 Thr Leu Leu Val Glu Ser Leu Ser Gly Ser Gln Thr Thr Gln Glu Leu Glu Asn Val Gly Arg Ile Val Gln Val Leu Arg Leu Leu Arg Ala Leu Arg Met Leu Lys Leu Gly Arg His Ser Thr Gly Leu Arg Ser Leu Gly Met Thr Ile Thr Gln Cys Tyr Glu Glu Val Gly Leu Leu Leu Phe Leu Ser Val Gly Ile Ser Ile Phe Ser Thr Val Glu Tyr Phe Ala Glu Gln Ser Ile Pro Asp Thr Thr Phe Thr Ser Val Pro Cys Ala Trp Trp Trp Ala Thr Thr Ser Met Thr Thr Val Gly Tyr Gly Asp Ile Arg Pro Asp Thr Thr Thr Gly Lys Ile Val Ala Phe Met Cys Ile Leu Ser Gly Ile Leu Val Leu Ala Leu Pro Ile Ala Ile Ile Asn Asp Arg Phe Ser

			420					425					130			
		125	Phe				440		Glu		- 4	145				
	AΕΛ	Leu				455	5		Ile	4	160					
4.00					470				Arg	475					400	
Arg	Leu	Lys	Gly	Arg		Arg	Ala	Ser	Thr 490	Arg :	Ser S	Ser (Gly (3ly . 495	qeA	
Asp	Phe	Trp		463												
		210>														
			342 DNA													
	<	213>	H.	sapi	.ens											
		220>														
			CDS (25		. (21	195)										
	<	223>	K+F	nov1	12											
	<	400>	9							2222	agat	+~~	നു ന	acco	ectect	. 60
	+	at a	2210	tccc	ccc a	agac	tcct	tc co	catct	cttt	agtt	CTTC	CE C	ccys	tteet	120
		+~+	ctar	racar		ccao	tttc	ct to	rttta	aca	qctc	:aagg	πgι	CLCC	aaguu	
ccca	cca	tcc	tgga	CC	ato (aca	σca	aac (ctq q	cc ac	g tg	g cu	g cc		ctggg t gct	
CCCS	,555		3 3-	1	Met .	Ala	Ala	Gly :	Leu A 5	la Th	r Tr	p Le	eu Pr	o Pl	ne Ala	ı
					1											
																240
cgg	gca	gca	gc	gt.	g gg	c to v Ti	g ct	g cc u Pr	c ccg	gcc Ala	cag Gln	caa Gln	ccc Pro	ctg Leu	ccc Pro	340
cgg Arg	gca	gca Ala 1	a Al	a gt	g gg 1 Gl	c to y Ti	p Le	g cc u Pr	c ccg o Pro	gcc Ala	cag Gln	caa Gln 25	ccc Pro	ctg Leu	Pro	340
Arg	Ala	Ala 1	A Al	a Va	l Gl	y Ti	p Le 2 :a to	u Pr 0	o Pro a qqa	gat	grg	25 gtt	ctg	gtg	gtg	340
Arg	gca Ala	Ala 1! a cc	A Al	a Va	l Gl	y Ti g go	np Le 2 ca to la Se	u Pr 0	c ccg o Pro a gga g Gly	gat	grg Xaa	25 gtt	ctg	gtg	gtg	
ecg Pro	gca Ala	Ala 1! a cc: a Pro	9 99 5 Gl	a Va g gt y Va	l Gl g aa l Ly	g ge	to Le ca to la Se 35	eu Pr :0 :t cg :r Ar	o Pro a gga g Gly	gat Asp	grg Xaa 40	25 gtt Val	ctg Leu	gtg Val	gtg Val	388
ccg Pro	gca Ala 30	Ala 1! a cc; a Pro	99 G1	a Va g gt y Va	g aa l Ly	g gg	to Lea to la Se	t cg	a gga g Gly	gat Asp	grg Xaa 40	25 gtt Val	ctg Leu	gtg Val	gtg Val	
ccg Pro	gca Ala 30 gtg Val	Ala 1! a cc; a Pro	99 G1	a Va g gt y Va	g aa l Ly g cg	g gg	to Lea to la Se	t cg	o Pro a gga g Gly	gat Asp	grg Xaa 40	25 gtt Val	ctg Leu	gtg Val	gtg Val	388
ecg Pro aac Asn 45	gea Ala 30 gtg Val	a Ala 1! a cc: a Pri	g gg c gg r Gl	g gt y Va a cg y Ar	g aa l Ly g cg g Ar	g gg s A	ta to la Se si la Se	et cg er Ar	a gga g Gly t tgg r Trp	gat Asp aag Lys 55	grg Xaa 40 aat Asn	gtt Val acg Thr	ctg Leu ctg Leu	gtg Val gac Asp	gtg Val cgc Arg 60	388
ecg Pro aac Asn 45	gea Ala 30 gtg Val	a Ala 1! a cc: a Pri	g gg c gg r Gl	g gt y Va a cg y Ar c tt	g aa l Ly g cg g Ar sg ct	g gg s A	ta to la Se si la Se	et cg er Ar	a gga g Gly t tgg r Trp	gat Asp aag Lys 55 aag	grg Xaa 40 aat Asn	gtt Val acg Thr	ctg Leu ctg Leu	gtg Val gac Asp	gtg Val cgc Arg 60	388 436
acc Pro aacc Asn 45 tacc	gca Ala 30 gts Val	a Ala 1! a cci a Pro) g ag l Se a ga o As	g gg c gg r Gl	g gt y Va a cg y Ar c tt	g aa l Ly g cg g Ar sg ct	g gg s A c t g P so sg g	ca to la Se la Se lt ga he Gl	et cg er Ar ag ac tu Th	a gga g Gly t tgg r Trr	gat Asp aag Lys 55 aag	grg Xaa 40 aat Asn gaa Glu	gtt Val acg Thr	ctg Leu ctg Leu ttc	gtg Val gac Asp tac Tyr	gtg Val cgc Arg 60 gat Asp	388 436 484
ccg Pro aac Asn 45 tac	gca Ala 30 gtc Val	Ala Com A Pro B aga B Se	g gg c gg r Gl	g gt y Va a cg y Ar c tt	g aa l Ly g cg g Ar sg ct	g ges Al	ca to la Se la Se la Se la Se lt ga lt ga lt Se	et cg er Ar ag ac lu Th	a gga g Gly t tgg r Trp g gag r Gli	gat Asp Asp Lys 55 aag Lys	grg Xaa 40 aat Asn gaa Glu	gtt Val acg Thr ttc Phe	ctg Leu ctg Leu ttc Phe	gtg Val gac Asp tac Tyr	gtg Val cgc Arg 60 gat Asp	388 436
ccg Pro aac Asn 45 tac	gca Ala 30 gtc Val	Ala Com A Pro B aga B Se	a Al gg gg gg gg gg Th	g gt y Va a cg y Ar c tt	g aa l Ly g cg g Ar sg ct	g ges Al	ca to la Se la Se la Se la Se lt ga lt ga lt Se	et cg er Ar ag ac lu Th gc to er Se tc ga	a gga g Gly t tgg r Trr	gat Asp Asp Lys 55 aag Lys	grg Xaa 40 aat Asn gaa Glu	gtt Val acg Thr ttc Phe	ctg Leu ctg Leu ttc Phe	gtg Val gac Asp tac Tyr 75	gtg Val cgc Arg 60 gat Asp	388 436 484
acc Pro acc Asn 45 tac Tyr	gca Ala 30 ya: Va:	1 Ala 1! 1 Cog Pri ag	a Al gg	y Va a cg y Ar c tt r Le c ga y GJ	g aa la Ly g cg cg Ar 5 5 cg ct Le Le C ta Le	g gg gg s Al c t t g g gg gg G gg g G G gg G G G gg G	rp Le 2 ca to ca 1 ca 5	nu Pr 00 ct cg car Ar ag acc lu Th ct cg car Ar	a gga g Gly t tgg r Trp g gag r Gli 70	gat gat aag bys 55 aag bys 55 aag bys 55 aag bys ctg	grg Xaa 40 aat Asn gaa Glu	gin 25 gtt Val acg Thr ttc Phe	ctg Leu ctg Leu ttc Phe atg	gtg Val gac Asp tac Tyr 75	gtg Val cgc Arg 60 gat Asp	388 436 484
acc Pro acc Asn 45 tac Tyr	gca Ala 30 ya: Va:	Ala 1! 1 con Private	a Al 99 Gl c gg Gl c ac ac ac ac ac ac ac gg Th a gg aa ac	y Va a cg y Ar c tt r Le c ga y GJ	g aa la Ly g cg cg Ar 5 5 cg ct Le Le C ta Le	g gg gg s Al c t t g g gg gg G gg g G G gg G G G gg G	TP Le 2 Ca tcc tc ggc agc agc agc agc agr Tr gga a arg T	tt cg gg gg hr G	a gga g Gly t tgg r Trp g gag r Gli 70	gat gat aag bys 55 aag bys 55 aag bys 55 aag bys ctg	grg Xaa 40 aat Asn gaa Glu	gtt Val acg Thr ttc Phe gac Asp	ctg Leu ctg Leu ttc Phe atg	gtg Val gac Asp tac Tyr 75	gtg Val cgc Arg 60 gat Asp	388 436 484 532
acc Asn 45 tac Tyr gct Ala	gca Ala 30 gto Va. Pro	a Ala 11 con a con	a Al gg gg Gl c gg gg aa gg aa gg aa agg ag a	a Va g gt y Va a cg c tt c ga cy Gl c tt n Pl	g aaal Ly g cg Ar s g ct s g ct tag ta	g gggs A. : ct gg P. : cg P. : cg P. : cg G. :	ca to la See la	t cg ac to ge to g	a gga g Gly t tgg r Trp g gag r Gli t cgg ap Arg	gat Asp Lys 55 aag Lys 25 gac 1 Asp 1 Ctg 1 Ctg 1 Ctg 1 Leu	grg Xaa 40 aat Asn gaa Glu cct Pro	gtt Val acg Thr ttc Phe gac Cys 105	ctg Leu ctg Leu ttc Phe atg	gtg Val gac Asp tac Tyr 75	gtg Val cgc Arg 60 gat Asp cgc Arg	388 436 484 532
aac Asn 45 tac Tyr get Ala	gca Ala 30 gtc Va. Pro	a Ala Con As	a Al gg gg gg gg gg gg gg aa gg aa gg aa gg aa aa	a Va ggt y Va a cg c tt r Le c ga y GJ c tt m Pl	g aaa ly g cg ctau le so tau le so t	g gggs A.: gg Pl gg G gg G gg G gg G tc tr	ca tcc a tcc	t cg ac to ga ac to g	a gga g Gly t tgg r Trp g gag r Gli 70 at egg p Arg	gat Asp SS	grg Xaaa 40 aat Asn gaa Glu cct Pro	gtt Val acg Thr ttc Phe gac Cys 105	ctg Leu ctg Leu ttc Phe atg	gtg Val gac Asp tac Tyr 75 ttc	gtg Val cgc Arg 60 gat Asp cgc Arg	388 436 484 532
aac Asn 45 tac Tyr get Ala	gca Ala 30 gtc Va. Pro	a Ala Con a	a Al gg gg gg gg gg gg gg aa gg aa gg aa gg aa aa	a Va ggt y Va a cg c tt r Le c ga y GJ c tt m Pl	g aaa ly g cg ctau le so tau le so t	y Tr g gg g gg g gg g g g g g g g g g ac t p p ac t p p ac t p p ac t p p	ca tcc a tcc	t cg ac to ga ac to g	a gga g Gly t tgg r Trp g gag r Gli t cgg ap Arg	gat Asp SS	grg Xaaa 40 aat Asn gaa Glu cct Pro	gtt Val acg Thr ttc Phe gac Cys 105	ctg Leu ctg Leu ttc Phe atg	gtg Val gac Asp tac Tyr 75 ttc	gtg Val cgc Arg 60 gat Asp cgc Arg	388 436 484 532
acc Asn 45 tac Tyr gct Ala cat His	gca Ala 30 gtc Val cca Pro ga As Val cca Pro 1 As	a Ala III Ala Company agents a gas a	a Al ggggggggggggggggggggggggggggggggggg	y Va cy Va cy Ar C cy G lice the Pl	g aaa l Ly g cg cg Ar s g ct s sg ta t	y Try g g g g g g g g g g g g g g g g g g g	ca to la Se	the Asset of Grand	a ggaggggggggggggggggggggggggggggggggg	gat Asp Lys 55 aagg Lys c gac Asp Asp Lys Asp Asp Asp Asp Asp Asp Asp Asp Asp As	grg Xaa 40 aat Asn gaa Glu cct Pro	acg Thr ttc Phe gac Cys 105 tacy 105	ctg Leu ctg Leu ttc Phe atg Met 90 cca Fro	gtg Val gac Asp tac Tyr 75 ttc Phe	gtg Val cgc Arg 60 gat Asp cgc Arg	388 436 484 532

130 135 140	
125 130 and gat gag gag gag gag gag ggg	724
aag gag aat gcc gag cgc ctg gca gag gat gag gag gab glu Glu Gln Ala Lys Glu Asn Ala Glu Arg Leu Ala Glu Asp Glu Glu Ala Glu Gln Ala 145 150 155	
ggg gac ggc cca gcc ctg cca gca ggc agc tcc ctg cgg cag cgg ctc Gly Asp Gly Pro Ala Leu Pro Ala Gly Ser Ser Leu Arg Gln Arg Leu 160 165 170	772
tgg cgg gcc ttc gag aat cca cac acg agc acc gca gcc ctc gtt ttc Trp Arg Ala Phe Glu Asn Pro His Thr Ser Thr Ala Ala Leu Val Phe 175 180 185	820
tac tat gtg acc ggc ttc ttc atc gcc gtg tcg gtc atc gcc aat gtg Tyr Tyr Val Thr Gly Phe Phe Ile Ala Val Ser Val Ile Ala Asn Val 190 195 200	868
gtg gag acc atc cca tgc cgc ggc tct gca cgc agg tcc tca agg gag Val Glu Thr Ile Pro Cys Arg Gly Ser Ala Arg Arg Ser Ser Arg Glu 205 210 215 220	916
cag ccc tgt ggc gaa cgc ttc cca cag gcc ttt ttc tgc atg gac aca Gln Pro Cys Gly Glu Arg Phe Pro Gln Ala Phe Phe Cys Met Asp Thr 225 230 235	964
gcc tgt gta ctc ata ttc aca ggt gaa tac ctc ctg cgg ctg ttt gcc Ala Cys Val Leu Ile Phe Thr Gly Glu Tyr Leu Leu Arg Leu Phe Ala 240 245 250	1012
gcc_ccc agc cgt tgc cgc ttc ctg cgg agt gtc atg agc ctc atc gac Ala Pro Ser Arg Cys Arg Phe Leu Arg Ser Val Met Ser Leu Ile Asp 255 260 265	1060
gtg gtg gcc atc ctg ccc tac tac att ggg ctt ttg gtg ccc aag aac Val Val Ala Ile Leu Pro Tyr Tyr Ile Gly Leu Leu Val Pro Lys Asn 270 275 280	1108
gac gat gtc tct ggc gcc ttt gtc acc ctg cgt gtg ttc cgg gtg ttt Asp Asp Val Ser Gly Ala Phe Val Thr Leu Arg Val Phe Arg Val Phe 295 300	1156
cgc atc ttc aag ttc tcc agg cac tca cag ggc ttg agg att ctg ggc Arg Ile Phe Lys Phe Ser Arg His Ser Gln Gly Leu Arg Ile Leu Gly 305 310	1204
tac aca ctc aag agc tgt gcc tct gag ctg ggc ttt ctc ctc ttt tcc Tyr Thr Leu Lys Ser Cys Ala Ser Glu Leu Gly Phe Leu Leu Phe Ser 320 325 330	1252
cta acc atg gcc atc atc atc ttt gcc act gtc atg ttt tat gct gag Leu Thr Met Ala Ile Ile Ile Phe Ala Thr Val Met Phe Tyr Ala Glu 335 340 345	1300
aag ggc aca aac aag acc aac ttt aca agc atc cct gcg gcc ttc tgg Lys Gly Thr Asn Lys Thr Asn Phe Thr Ser Ile Pro Ala Ala Phe Trp 350 355 360	1348
tat acc att gtc acc atg acc acg ctt ggc tac gga gac atg gtg ccc Tyr Thr Ile Val Thr Met Thr Thr Leu Gly Tyr Gly Asp Met Val Pro 365 370 380	1396

agc acc att gct ggc aag att ttc ggg tcc atc tgc tca ctc agt ggc Ser Thr Ile Ala Gly Lys Ile Phe Gly Ser Ile Cys Ser Leu Ser Gly 385 390 395	1444
gtc ttg gtc att gcc ctg cct gtg cca gtc att gtg tcc aac ttt agc Val Leu Val Ile Ala Leu Pro Val Pro Val Ile Val Ser Asn Phe Ser 400 405 410	1492
cgc atc tac cac cag aac cag cgg gct gac aag cgc cga gca cag cag Arg Ile Tyr His Gln Asn Gln Arg Ala Asp Lys Arg Arg Ala Gln Gln 415 420 425	1540
aag gtg cgc ttg gca agg atc cga ttg gca aag agt ggt acc acc aat Lys Val Arg Leu Ala Arg Ile Arg Leu Ala Lys Ser Gly Thr Thr Asn 430 435 440	1588
gcc ttc ctg cag tac aag cag aat ggg ggc ctt gag gac agc ggc agt Ala Phe Leu Gln Tyr Lys Gln Asn Gly Gly Leu Glu Asp Ser Gly Ser 450 455 460	1636
ggc gag gaa cag gct ctt tgt gtc agg aac cgt tct gcc ttt gaa cag Gly Glu Glu Gln Ala Leu Cys Val Arg Asn Arg Ser Ala Phe Glu Gln 465 470 475	1684
caa cat cac cac ttg ctg cac tgt cta gag aag aca acg tgc cat gag Gln His His Leu Leu His Cys Leu Glu Lys Thr Thr Cys His Glu 480 485 490	1732
ttc aca gat gag ctc acc ttc agt gaa gcc ctg gga gcc gtc tcg ccg Phe Thr Asp Glu Leu Thr Phe Ser Glu Ala Leu Gly Ala Val Ser Pro 495 500 505	1780
ggt ggc cgc acc agc cgt agc acc tct gtg tct tcc cag cca gtg gga Gly Gly Arg Thr Ser Arg Ser Thr Ser Val Ser Ser Gln Pro Val Gly 510 515	1828
ccc gga agc ctg ctg tct tct tgc tgc cct cgc agg gcc aag cgc cgc Pro Gly Ser Leu Leu Ser Ser Cys Cys Pro Arg Arg Ala Lys Arg Arg 525 530 535	1876
gcc atc cgc ctt gcc aac tcc act gcc tca gtc agc cgt ggc agc atg Ala Ile Arg Leu Ala Asn Ser Thr Ala Ser Val Ser Arg Gly Ser Met 545 550 555	1924
cag gag ctg gac atg ctg gca ggg ctg cgc agg agc cat gcc cct cag Gln Glu Leu Asp Met Leu Ala Gly Leu Arg Arg Ser His Ala Pro Gln 560 565 570	1972
age ege tee age ete aat gee aag eee eat gae age ett gae etg aac Ser Arg Ser Ser Leu Asn Ala Lys Pro His Asp Ser Leu Asp Leu Asn 575 580 585	2020
tgc gac agc cgg gac ttc gtg gct gcc att atc agc atc cct acc cct Cys Asp Ser Arg Asp Phe Val Ala Ala Ile Ile Ser Ile Pro Thr Pro 590 595 600	2068
cct gcc aac acc cca gat gag agc caa cct tcc tcc cct ggc ggc ggt Pro Ala Asn Thr Pro Asp Glu Ser Gln Pro Ser Ser Pro Gly Gly 605 610 615 620	2116

```
ggc agg gcc ggc agc acc ctc agg aac tcc agc ctg ggt acc cct tgc
Gly Arg Ala Gly Ser Thr Leu Arg Asn Ser Ser Leu Gly Thr Pro Cys
                                  630
               625
                                                                  2215
ctc ttc ccc gag act gtc aag atc tca tcc c tgtgaggggt aggcctgctg
Leu Phe Pro Glu Thr Val Lys Ile Ser Ser
           640
attcagaggg teetetteat ttttgggaae teettteeaa agecatattt ttgggaggea
                                                                  2275
gagagggca ggcttgggca ccccttctgc ccccccact gagaactatg caatggagtt
                                                                  2335
tcatgaaatg gtccacatag tggggaagta gccaggaaat gagaaacttc ctcccacccc
                                                                  2395
agacattttt cctggtggga gctgaagcac tgggcttcca caggcccctg gcctccttgc
                                                                  2455
cctagcacac tgggactggc cccactctcc cagctggact cctgcatgct cctccccttg
                                                                   2515
ggeteteaga tgaaggeaaa getttgatee gacatetgag etetageeta agaaggagag
                                                                   2575
ttgagatttc etectecete tggetgggat atggagettt ggaggttcag agaagagaac
                                                                   2635
cetcacetet gatetggeet ctacgagagg teetcatete catetggeee aacaatteee
                                                                   2695
agattetgaa gettggaatg caaacacagg etteatggge tgtggeetet geagegaeet
gccatcccca ggccttgcct gaggggtcag gctgcctctc ccaacacaca ctcagatagc
                                                                   2815
acaaattcta ccatcccctt ccctggctgc tggaaatgga ccccgcaacc ctgtcctctg
                                                                   2875
etgggeecee agcaaactet agcaatagca getgetgeeg tgteattatg caaageetet
                                                                   2935
gaccagtttg etgeageatt tacatetgee etaateagag gggeeacete taaeteetee
                                                                   2995
tectectete ttetectetg gtttgegtee tteetgggtt gggetggagt etggaetgge
tgagataaga gcctggcaac cagcaagagc tgggctgtat ttggagatca tgggctgatt
                                                                   3115
ccatgttctt gggcaacagt ccagaagcat caggggctcc ggcctgggat gtttctgaac
                                                                   3175
tttgggagtt ataggagaca ggaggaactt etecteetee teeteecta caatteettt
                                                                   3235
tcacatattc ctttcttctc cctcttgggt gaccttccaa aactctgctc tcaggctgaa
atctggcatc atctcaggtt ccctgtcccc agcactgtcc ccatggagct ggtggctgac
                                                                   3355
 3415
                                                                   3424
 aaaaaaaaa
       <210> 10
       <211> 646
       <212> PRT
     <213> H. sapiens
       <220>
       <221> VARIANT
       <222> (1)...(646)
       <223> Xaa = Any Amino Acid
 Met Ala Ala Gly Leu Ala Thr Trp Leu Pro Phe Ala Arg Ala Ala Ala
                                   10
                  5
 Val Gly Trp Leu Pro Pro Ala Gln Gln Pro Leu Pro Pro Ala Pro Gly
                                 25
  Val Lys Ala Ser Arg Gly Asp Xaa Val Leu Val Val Asn Val Ser Gly
             20
                             40
  Arg Arg Phe Glu Thr Trp Lys Asn Thr Leu Asp Arg Tyr Pro Asp Thr
                                            60
                         55
  Leu Leu Gly Ser Ser Glu Lys Glu Phe Phe Tyr Asp Ala Asp Ser Gly
                                         75
                     70
  Glu Tyr Phe Phe Asp Arg Asp Pro Asp Met Phe Arg His Val Leu Asn
                                    90
  Phe Tyr Arg Thr Gly Arg Leu His Cys Pro Arg Gln Glu Cys Ile Gln
                 85
                                105
  Ala Phe Asp Glu Glu Leu Ala Phe Tyr Gly Leu Val Pro Glu Leu Val
                             120
  Gly Asp Cys Cys Leu Glu Glu Tyr Arg Asp Arg Lys Lys Glu Asn Ala
                                             140
                         135
  Glu Arg Leu Ala Glu Asp Glu Glu Ala Glu Gln Ala Gly Asp Gly Pro
                                          155
                      150
```

Ala Leu Pro Ala Gly Ser Ser Leu Arg Gln Arg Leu Trp Arg Ala Phe 175 170 185 185 185 186 187 188 188 188 188 188 188 188 188 188
Ala Leu Pro Ala Gly Ser Ser Leu Als 170
The Ser Thr Ala Ala Leu val File 190
Glu Asn Pro His Thr Ser Val 185 180 185 186 Gly Phe Phe Ile Ala Val Ser Val Ile Ala Asn Val Val Glu Thr Ile 205 200 200 200 200 200 200 20
Cly phe Phe Ile Ala Val Ser Val Ile Ala Assa 205
Gly Phe Phe Ite Ala Value 200 195 Pro Cys Arg Gly Ser Ala Arg Arg Ser Ser Arg Glu Gln Pro Cys Gly 220 220 220 220 220 220 220 2
Pro Cys Arg Gly Ser Ala Arg Arg 220
210 240
Glu Arg Phe Pro Girl 230 235 Phe Ala Ala Pro Ser Arg
Glu Arg Phe Pro Gli 230 230 230 225 11e Phe Thr Gly Glu Tyr Leu Leu Arg Leu Phe Ala Ala Pro Ser Arg 255 11e Phe Thr Gly Glu Tyr Leu Leu Arg Leu Phe Ala Ala Pro Ser Arg 255 11e Phe Thr Gly Glu Tyr Leu Leu Arg Leu Phe Ala Ala Pro Ser Arg 255 11e Phe Thr Gly Glu Tyr Leu Leu Arg Leu Phe Ala Ala Pro Ser Arg 255 11e Phe Thr Gly Glu Tyr Leu Leu Arg Leu Phe Ala Ala Pro Ser Arg 255
245 Net Ser Leu Ile Asp Val Ala Ile
Ile Phe Thr Gly Glu 172 250 250 250 245 270 265 265 250 Asp
The Gly Leu Leu Val Pro Lys Asia 125
Leu Pro Tyr Tyr IIe 617 280 280 275 280 Ala Phe Arg Ile Phe Lys 300 300 295 295 200 Cly Tyr Thr Leu Lys
Gly Ala Phe Val Thr Leu Arg Val Phe Arg 300
Gly Ala Phe Val Thr 295 290 290 Phe Ser Arg His Ser Gln Gly Leu Arg Ile Leu Gly Tyr Thr Leu Lys 310 320 Phe Ser Arg His Ser Gln Gly Leu Arg Ile Leu Gly Tyr Thr Leu Lys 320 Phe Ser Leu Thr Met Ala
Phe Ser Arg His Ser Gill Gry 200 315
305 Gly Ley Gly Phe Leu Leu Phe Ser 235
Ser Cys Ala Ser Giu Lea 37 330 325 Ile Ile Ile Phe Ala Thr Val Met Phe Tyr Ala Glu Lys Gly Thr Asn 350 345 345 347 348 349 340 340
The Ile Ile Phe Ala Thr Val Met Phe 191 350
The Cor Tie Pro Ala Ala Phe 119
Lys Thr Asn Phe Thr Sel 360 355 Thr Met Thr Thr Leu Gly Tyr Gly Asp Met Val Pro Ser Thr Ile Ala 380 375 The Met Thr Thr Leu Gly Tyr Gly Asp Met Val Leu Val Ile
355 The The Leu Gly Tyr Gly Asp Met Val Flo 380
Thr Met Thr Leu Gly 17-380 375 375 370 Gly Lys Ile Phe Gly Ser Ile Cys Ser Leu Ser Gly Val Leu Val Ile 400 Gly Lys Ile Phe Gly Ser Ile Cys Ser Leu Ser Arg Ile Tyr His
Gly Lys Ile Phe Gly Ser He Cys Box 395
385 val Tie Val Ser Ash Fite 3415
Ala Leu Pro Val Pro Val 200 410 405 Gin Asn Gln Arg Ala Asp Lys Arg Arg Ala Gln Gln Lys Val Arg Leu 430 425 The Arg Ala Phe Leu Gln
Gir Asn Gln Arg Ala Asp Lys Arg Arg Ala Gir 430
420 Ala Iva Ser Gly Thr Thr Asn Ala Phe Leu Gin
Gln Asn Gln Arg Ala Asp 272 425 420 Ala Arg Ile Arg Leu Ala Lys Ser Gly Thr Thr Asn Ala Phe Leu Gln 445 440 Ala Arg Gly Ser Gly Glu Glu Gln
435 Gly Gly Leu Glu Asp Ser Gly 365
Tyr Lys Gln Asn Gly Gly 455 450 450 Ala Leu Cys Val Arg Asn Arg Ser Ala Phe Glu Gln Gln His His His 480 470 470 470 470 470 480 470 470 470 470 470 470 470 470 470 47
Ala Leu Cys Val Arg Asn Arg Sel Ala 475
465 The Clu Lys The The Cys His Glu 1495
Leu Leu His Cys Leu Glu Ala Leu Gly Ala Val Ser Pro Gly Gly Arg Thr Leu Thr Phe Ser Glu Ala Leu Gly Ala Val Ser Pro Gly Ser Leu 505 505 505 507 508 509 509 500
Tou Thr Phe Ser Glu Ala Leu Gly Ala Val Ser 130 510
500 Ser Gln Pro Val Gly Pro Gly Ser Leu
Ser Arg Ser Thr Ser Val Ser Ser Gln Pro Val Gly Pro Gly Ser Leu Ser Arg Ser Thr Ser Val Ser Ser Gln Pro Val Gly Pro Gly Ser Leu Ser Arg Ser Thr Ser Val Ser Ser Gln Pro Val Gly Pro Gly Ser Leu Ser Arg Ser Thr Ser Val Ser Ser Gln Pro Val Gly Pro Gly Ser Leu
515 Gree Pro Arg Arg Ala Lys Arg Arg
Leu Ser Ser Cys Cys To 535 530 Ala Asn Ser Thr Ala Ser Val Ser Arg Gly Ser Met Gln Glu Leu Asp 550 550 550 560 570 580 580 580 580 580 580 58
Ala Asn Ser Thr Ala Ser Val Ser Aig 617 555
545 Ser Arg Ser His Ala Pro Gln Ser Arg Ser
Ala Asn Ser Thr Ala 555 555 555 556 557
Dro His Asp Ser Leu Asp Leu Ash 590
Leu Asn Ala Lys Flo Lie 585 580 Asp Phe Val Ala Ala Ile Ile Ser Ile Pro Thr Pro Pro Ala Asn Thr 605 600 600 601 Gly Gly Arg Ala Gly
Asp Phe Val Ala Ala Ile IIe Ser IIe 120 605
Sys Cla Pro Ser Ser Pro Gly Gly Gly
Pro Asp Glu Ser Gin Fig. 615 620 Glu Phe Pro Glu
Pro Asp Glu Ser Gli Fin 615 610 610 Ser Thr Leu Arg Asn Ser Ser Leu Gly Thr Pro Cys Leu Phe Pro Glu 640 630
mber Val Lvs Ile Ser Ser
7mi vaz -1 20

<210> 11	
<211> 1862	
<212> DNA	
<213> H. sapiens	
<220>	
<221> CDS	
<222> (383)(1157)	
<223> K+Hnov15	
(2207	
<400> 11	60
<400> 11 cagctgaatg tggaggcctt taagagaact tccagctcct gtaaaaaccc agaccagagg	120
cagetgaatg tggaggeett taagagaaet tecageteet gedadate cettaetee actaetgace aacattteag getgateete cagacetega agttaetete ettaetetee actaetgace aacattteag getgateete gaacatetegg gaaaagaetg aagaaataat	180
actactgacc aacatttcag gctgatcctc cagacttcga agetacagactg aagaaataat tgactcttaa ttacatcaca cctgtgtcga cactctctgg gaaaagactg aagaaataat tgactcttaa ttacatcaca actgtgtcga ataggctgat acgccaccta ctgcaaaacc	240
tgactettaa ttacateaca eetgtgtega caetetetgg gaddaga-tgategategategatac etgeaaaace etttteaaga agcagaaage teetgeatac ataggetgat aegeeaceta etgeaaaace etttteaaga agcagaaage teetgeatac ateaceagge tggggetgaa	300
cttttcaaga agcagaaagc tcctgcatac ataggctgat acgcacaggc tgggggctgaa gagctgacag cgcaggcgat gctgccagcg tttccattcc atcaccaggc tggggctgaa gagctgacag cgcaggcgat gctgccagcg tttcaaaaaaa ctcaaagcca agaagaacaa	360
gagetgacag egeaggegat getgecageg tttecattet attaceaggea agaagaacaa taaaggegtg ettgtgtggt agtgtetett tttaaaaaaat etcaaagcea agaagaacaa taaaggegtg ettgtgtggt agtgtetett tttaaaaaaat etcaaagcea agaagaacaa	412
taaaggcgtg cttgtgtggt agtgtctctt tttaaaaaat ctcaaaga aga gaa aaa gctgaaatag catcttcaaa aa atg gag cgt aaa ata aac aga aga gaa aaa gctgaaatag catcttcaaa aa atg gag cgt aaa ata aac aga aga gaa aaa	
10	
1 5	
got act gat caa	460
gaa aag gag tat gaa ggg aaa cac aac agc ctg gaa gat act gat caa	
al., Ive Clu TVr Glu Gly Lys Mas 25	
15	
at and get age tat	508
gga aag aac tgc aaa tcc aca ctg atg acc ctc aac gtt ggt gga tat	
Cly Lys Asn Cys Lys Ser IIII Dea 110	
30	
tag coa gag act tto	556
tta tac att act caa aaa caa aca ctg acc aag tac cca gac act ttc	
Len Tvr Ile Thr Gin Lys Gin im 255	
45	
ctt gaa ggt ata gta aat gga aaa atc ctc tgc ccg ttt gat gct gat	604
ctt gaa ggt ata gta aat gga aaa atc tte tys pro Phe Asp Ala Asp	
Leu Glu Gly Ile Val Ash Gry 170	
60	
ggt cat tat ttc ata gac agg gat ggt ctc ctc ttc agg cat gtc cta	652
ggt cat tat ttc ata gac agg gat ggt tou Leu Phe Arg His Val Leu	
Gly His Tyr Phe He Asp Ang Ang 527	
75	
aac ttc cta cga aat gga gaa ctt cta ttg ccc gaa ggg ttt cga gaa	700
aac ttc cta cga aat gga gaa ctt ctu bos Pro Glu Gly Phe Arg Glu	
Asn Phe Leu Arg Ash Gly Glu Leu 100 105	
95	740
aat caa ctt ctt gca caa gaa gca gaa ttc ttt cag ctc aag gga ctg	748
aat caa ctt ctt gca caa gda gca gdu Phe Phe Gln Leu Lys Gly Leu	
Asn Gln Leu Leu Ala Gin Glu Ala 115 120	
110	700
gca gag gaa gtg aaa too agg tgg gag aaa gaa cag ota aca coo aga	796
gca gag gaa gtg aaa too ayy typ Glu Lys Glu Gln Leu Thr Pro Arg	
Ala Glu Glu Val Lys Sel Alg 120 135	
125	044
gag act act ttc ttg gaa ata aca gat aac cac gat cgt tca caa gga	844
gag act act ttc ttg gaa ata aca gat aac cac gat ogs of Gly Glu Thr Thr Phe Leu Glu Ile Thr Asp Asn His Asp Arg Ser Gln Gly	
Glu Thr Thr Phe Leu Glu 11e 111 ASP ASIA 150	
140	002
tta aga atc ttc tgt aat gct cct gat ttc ata tca aaa ata aag tct	892
tta aga atc ttc tgt aat gct cct gat ttc ata tta dda dte Lys Ser Leu Arg Ile Phe Cys Asn Ala Pro Asp Phe Ile Ser Lys Ile Lys Ser	
Leu Arg Ile Phe Cys Ash Ala Flo Asp 120	

165	170
155	940
cgc att gtt ctg gtg tcc aaa agc agg ctg gat g Arg Ile Val Leu Val Ser Lys Ser Arg Leu Asp 175	raa ftt cca yay yay
ttt tca ata tcg tca aat atc atc caa ttt aaa Phe Ser Ile Ser Ser Asn Ile Ile Gln Phe Lys 190 195	tac ttc ata aag tct 988 Tyr Phe Ile Lys Ser 200
gaa aat ggc act cga ctt gta cta aag gaa gac Glu Asn Gly Thr Arg Leu Val Leu Lys Glu Asp 205 210	215
acc ttg gaa act ctt aag ttt gag gct atc atg Thr Leu Glu Thr Leu Lys Phe Glu Ala Ile Met 220 225	atg gct tta aag tgt 1084 Met Ala Leu Lys Cys 230
ggc ttt aga ctg ctg acc agc ctg gat tgt tcc Gly Phe Arg Leu Leu Thr Ser Leu Asp Cys Ser	aaa ggg tca att gtt 1132 Lys Gly Ser Ile Val 250
cac agc gat gca ctt cat ttt atc a agtaattac His Ser Asp Ala Leu His Phe Ile	c tgtgtcacga 1177
255	and apparent 1237
acaaaggcaa caagcatgca gccagcaagc ttcggaaaac	caracted dayacaroo
anataacatg cccagctage telglaceae assist	- staggatatt cctactgatc 1357
ctaacggtat gtaaattcta tcgctaaaga ago	actgtaaata aagactgaaa 141/
agactettee acctaaaatg aaaacagtaa cettetata gettttgeta tttatttgte ettaagetgt ettteaatt	agattgtctt gggtatttgc 14//
gettttgeta tttatttgte ettaagetgt ettedatt acaaaaagaa geatgtacat tatetategt teatttaag	t aaatggtaat aaaatattt 1557
aaggggctat taatatttaa aattettaa	a atchactaa aacagagcta 1657
actogcaaaa ttaactacct ggagcaaaac agassa	a troatttgat ttttccatag 1/1/
tagtgaaaca aaatgagatt gtaagaaga	+ arragactta tagctgaatt 1///
ggtatttac tqaaaattcc tagaaattcc	a ataaaaagta aataaaagta 1862
ctgctacctt caaaaaaaaa aaaaa	
-	
<210> 12 <211> 258	
<212> PRT	
<213> H. sapiens	
<400> 12	a run Glu Tur Glu Gly
<400> 12 Met Glu Arg Lys Ile Asn Arg Arg Glu Lys G 10	15
1 5 10 Lys His Asn Ser Leu Glu Asp Thr Asp Gln G	ly Lys Asn Cys Lys Ser
Lys His Asn Ser Leu Glu 125	30
20 25 Thr Leu Met Thr Leu Asn Val Gly Gly Tyr L	eu Tyr 11e 111 Gin 27
35 40 Gln Thr Leu Thr Lys Tyr Pro Asp Thr Phe I	eu Glu Gly Ile Val Asn
Gln Thr Leu Thr Lys Tyr Flo Asp 1112 55	60
50 55 Gly Lys Ile Leu Cys Pro Phe Asp Ala Asp (Sly His Tyr Phe Tie Aby
65 70 Arg Asp Gly Leu Leu Phe Arg His Val Leu Phe Arg His Val Leu Phe Phe Arg His Val Leu Phe	Asn Phe Leu Arg Asn Gly
Arg Asp Gly Leu Leu Phe Arg His var het 2	95
85 90 Glu Leu Leu Pro Glu Gly Phe Arg Glu	Asn Gln Leu Leu Ala Gin
100 105 Glu Ala Glu Phe Phe Gln Leu Lys Gly Leu	Ala Glu Glu Val Lys Ser
Glu Ala Glu Phe Phe Gln Leu Lys Gly Beu 115	125

The Dro Arg Glu Thr Thr Phe Leu Glu	
Arg Trp Glu Lys Glu Gln Leu Thr Pro Arg Glu Thr Thr Phe Leu Glu	
135 130 135 140 130 135 140 140 140 150 150 150 150 150 150 150 150 150 15	
150 150 150 155 Leu Val Leu Val Ser	
150 145 150 150 150 155 145 Ala Pro Asp Phe Ile Ser Lys Ile Lys Ser Arg Ile Val Leu Val Ser 175 170 170 175 176 177 178 179 179 170 170 170 170 170 170 170 170 170 170	
and Arg Leu Asp Gly Phe Pro Glu Glu Phe Ser 11e Ser Ser 190	
Lys Ser Arg Leu 185 180 185 Asn Gly Thr Arg Leu	
185 180 180 181 182 185 186 187 188 189 180 180 181 185 205 200 200 200 200 200 200 200 200 20	
195 Val Leu Lys Glu Asp Asn Thr Phe Val Cys Thr Leu Glu Thr Leu Lys 220 215 220 215 220 215 220 217 220 218 220 219 220 210 210 220	
210 210 215 220 217 Phe Glu Ala Ile Met Met Ala Leu Lys Cys Gly Phe Arg Leu Leu Thr 240 235 230 235 237 242 Leu His	
230 230 His Ser Asp Ala Leu His	
230 225 Ser Leu Asp Cys Ser Lys Gly Ser Ile Val His Ser Asp Ala Leu His 250 255 265	
phe Ile	
<210> 13	
<211> 1877	
<212> DNA	
<213> H. sapiens	
.2205	
<220> <221> CDS	
<222> (322)(1090)	
<223> K+Hnov27	
100. 13	60
<400> 13 caccacegee eccageegee etegetgggg aacaettaca teeteeceaa agacageeag caccacegee eccageegee etegetgggg eccaagegeg eccetgtaega gtetgtgtte	120
	180 240
	300
	351
acccaggaca gtcggcccaa t atg tca aga cct ctg atc ass as acccaggaca gtcggcccaa t atg tca aga cct ctg atc ass as acccaggaca gtcggcccaa t atg tca aga cct ctg atc ass as acccaggaca gtcggcccaa t atg tca aga cct ctg atc ass as	
1	399
gca tot cca otg awo aac caa ggo ato cot act cca gca caa oto aca	300
gca tot oca otg awo aac caa ggc ato cot act coa gca cub leu Thr Ala Ser Pro Leu Xaa Asn Gln Gly Ile Pro Thr Pro Ala Gln Leu Thr 20	
aaa too aat gog cot gto cac att gat gtg ggc ggc cac atg tac acc	447
aaa too aat gog cot gto cac att gat gog 959 55 His Met Tyr Thr	
Lys Ser Asn Ala Flo Val	
30 aga atc qqa aga	495
age age ctg gee ace etc ace aaa tae ect gaa tee aga ate gga aga Ser Ser Leu Ala Thr Leu Thr Lys Tyr Pro Glu Ser Arg Ile Gly Arg 50	
Ser Ser Leu Ala Thr Leu Thr Lys Tyr Pro Glu 555	
Δ h	543
ctt ttt gat ggt aca gag ccc att gtt ttg gac agt ctc aaa cag cac	
ctt ttt gat ggt aca gag ccc att gtt ttg gac agt ttt dat 555 Leu Phe Asp Gly Thr Glu Pro Ile Val Leu Asp Ser Leu Lys Gln His 65 70	
	F01
the state of the sate till	591
tat ttc att gac aga gat gga cag des Phe Arg Tyr Ile Leu Ash Phe	
Tyr Phe lie ASP Ary 80	
75	639
cta cga aca tcc aaa ctc ctc att cct gat gat ttc aag gac tac act	
ten Ard Thr Ser bys 255 -	
23	

95 100 105	
ttg tta tat gaa gag gca aaa tat ttt cag ctt cag ccc atg ttg ttg Leu Leu Tyr Glu Glu Ala Lys Tyr Phe Gln Leu Gln Pro Met Leu Leu	687
gag atg gaa aga tgg aag cag gac aga gaa act ggt cga ttt tta agg Glu Met Glu Arg Trp Lys Gln Asp Arg Glu Thr Gly Arg Phe Ser Arg	735
ccc tgt gag tgc ctc gtc gtg cgt gtg gcc cca gac ctc gga gaa agg Pro Cys Glu Cys Leu Val Val Arg Val Ala Pro Asp Leu Gly Glu Arg 145	783
atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg ata gaa gaa gta ttt cca gag atc acg cta agc ggt gac aaa tcc ttg atc acg cta gac gac gac gac gac gac gac gac gac ga	831
atc ggc gac gtg atg tgt aac tct gtc aat gca ggc tgg aat cac gac atc ggc gac gtg atg tgt aac tct gtc aat gca ggc tgg aat cac gac atc ggc gac gtg atg tgt aac tct gtc aat gca ggc tgg aat cac gac atc ggc gac gtg atg tgt aac tct gtc aat gca ggc tgg aat cac gac atc ggc gac gtg atg tgt aac tct gtc aat gca ggc tgg aat cac gac atc ggc gac gtg atg tgt aac tct gtc aat gca ggc tgg aat cac gac atc ggc gac gtg atg tgt aac tct gtc aat gca ggc tgg aat cac gac atc ggc ggc tgg aat cac gac	879
tcg acg cac gtc atc agg ttt cca cta aat ggc tac tgt cac ctc aac Ser Thr His Val Ile Arg Phe Pro Leu Asn Gly Tyr Cys His Leu Asn 190 195	927
tca gtc cag gtc ctc gag agg ttg cag caa aga gga ttt gaa atc gtg Ser Val Gln Val Leu Glu Arg Leu Gln Gln Arg Gly Phe Glu Ile Val 205 210 215	975
ggc tcc tgt ggg gga gga gta gac tcg tcc cag ttc agc gaa tac gtc Gly Ser Cys Gly Gly Gly Val Asp Ser Ser Gln Phe Ser Glu Tyr Val 220 225 230	1023
ctt cgg cgg gaa ctg agg cgg acg ccc cgt gta ccc tcc gtc atc cgg Leu Arg Arg Glu Leu Arg Arg Thr Pro Arg Val Pro Ser Val Ile Arg 240 245	1071
ata aag caa gag cct ctg g actaaatgga catatttctt atgcaaaaag Ile Lys Gln Glu Pro Leu 255	1120
	1180
gaaaacacac acaaccaata actcaaacaa aaaagggaca tttatgtgca gttgggacag	1240
caaaccaaqt cctggacgta adattgdda hartatat caaggtgtaa aaaatatata	1300 1360
antatata tatatgtcaa aagguaggaa aagguaggaa ataaaaaaaa	1420
accordatto otoctotoat yyetysys s	1480
tagacaadcc atdagtggtg aggustus	1540 1600
nacaaaacac cttqaatcaa gutuguugu aana tagaaagatg ttagacatga	1660
gggaccaggc aggacttcag adadaccoot and garactcaca gatgtgaact	1/20
aattttaaat qtagtttgta tagaastaatta gaattattcc ttqttagaat	1,00
	1877
tgctccagtt caagtctgct goodbaaaaaaa aaaaaaaa caataaactc tgtttaaaaa ataaaaaaaaa aaaaaaaa	
<210> 14	
<211> 256 <212> PRT	
<212> PR1 <213> H. sapiens	

Add	<220> <221> VARIANT <222> (1)(256) <223> Xaa = Any Amino Acid	
His Ile Asp Val Gly Gly His Met Tyr Thr Ser Ser Leu Ala Thr Leu 35 40 Thr Lys Tyr Pro Glu Ser Arg Ile Gly Arg Leu Phe Asp Gly Thr Glu 50 Fro Ile Val Leu Asp Ser Leu Lys Gln His Tyr Phe Ile Asp Arg Asp 65 Fro Ile Val Leu Asp Ser Leu Lys Gln His Tyr Phe Ile Asp Arg Asp 66 670 680 Fro Ile Val Leu Asp Ser Leu Lys Gln His Tyr Phe Ile Asp Arg Asp 670 681 Eleu Ile Pro Asp Asp Phe Lys Asp Tyr Thr Leu Leu Tyr Glu Glu Ala 670 Lys Tyr Phe Gln Leu Gln Pro Met Leu Leu Glu Met Glu Arg Trp Lys 671 180 Lys Tyr Phe Gln Leu Gln Pro Met Leu Leu Glu Met Glu Arg Trp Lys 671 181 182 Cln Asp Arg Glu Thr Gly Arg Phe Ser Arg Pro Cys Glu Cys Leu Val 681 183 Val Arg Val Ala Pro Asp Leu Gly Glu Arg Ile Thr Leu Ser Gly Asp 685 Asn Ser Val Asn Ala Gly Trp Asn His Asp Ser Thr His Val Ile Arg 686 Phe Pro Leu Asn Gly Tyr Cys His Leu Asn Ser Val Gln Val Leu Glu 687 Arg Leu Gln Gln Arg Gly Phe Glu Tle Val Gly Ser Cys Gly Gly Gly 687 Arg Leu Gln Gln Arg Gly Phe Glu Tyr Val Leu Arg Arg Glu Leu Arg 687 2887 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 688 4210 4215 4210 4215 4210 4215 4210 4215 4210 4	<pre> <400> 14 Met Ser Arg Pro Leu Ile Thr Arg Ser Pro Ala Ser Pro Leu Xaa Asn 15 10 15</pre>	
His Ile Asp Val Gly Gly His Met Tyr Thr Ser Ser Leu Ala Thr Leu 35 40 Thr Lys Tyr Pro Glu Ser Arg Ile Gly Arg Leu Phe Asp Gly Thr Glu 50 Fro Ile Val Leu Asp Ser Leu Lys Gln His Tyr Phe Ile Asp Arg Asp 65 Fro Ile Val Leu Asp Ser Leu Lys Gln His Tyr Phe Ile Asp Arg Asp 66 670 680 Fro Ile Val Leu Asp Ser Leu Lys Gln His Tyr Phe Ile Asp Arg Asp 670 681 Eleu Ile Pro Asp Asp Phe Lys Asp Tyr Thr Leu Leu Tyr Glu Glu Ala 670 Lys Tyr Phe Gln Leu Gln Pro Met Leu Leu Glu Met Glu Arg Trp Lys 671 180 Lys Tyr Phe Gln Leu Gln Pro Met Leu Leu Glu Met Glu Arg Trp Lys 671 181 182 Cln Asp Arg Glu Thr Gly Arg Phe Ser Arg Pro Cys Glu Cys Leu Val 681 183 Val Arg Val Ala Pro Asp Leu Gly Glu Arg Ile Thr Leu Ser Gly Asp 685 Asn Ser Val Asn Ala Gly Trp Asn His Asp Ser Thr His Val Ile Arg 686 Phe Pro Leu Asn Gly Tyr Cys His Leu Asn Ser Val Gln Val Leu Glu 687 Arg Leu Gln Gln Arg Gly Phe Glu Tle Val Gly Ser Cys Gly Gly Gly 687 Arg Leu Gln Gln Arg Gly Phe Glu Tyr Val Leu Arg Arg Glu Leu Arg 687 2887 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 688 4210 4215 4210 4215 4210 4215 4210 4215 4210 4	Gln Gly Ile Pro Thr Pro Ala Gln Leu Thr Lys Ser Asn Ala Pro Val	
The Lys Tyr Pro Glu Ser Arg The Gly Arg Leu Phe Asp Gly Thr Glu 50 Fro Ile Val Leu Asp Ser Leu Lys Gln His Tyr Phe Ile Asp Arg Asp 61 Fro Ile Val Leu Asp Ser Leu Lys Gln His Tyr Phe Ile Asp Arg Asp 65 Fro Ile Val Leu Asp Ser Leu Lys Gln His Tyr Phe Ile Asp Arg Asp 65 Gly Gln Met Phe Arg Tyr Ile Leu Asn Phe Leu Arg Thr Ser Lys Leu 90 Leu Ile Pro Asp Asp Phe Lys Asp Tyr Thr Leu Leu Tyr Glu Glu Ala 100 Lys Tyr Phe Gln Leu Gln Pro Met Leu Leu Glu Met Glu Arg Trp Lys 115 110 Lys Tyr Phe Gln Leu Gln Pro Met Leu Leu Glu Met Glu Arg Trp Lys 115 Gln Asp Arg Glu Thr Gly Arg Phe Ser Arg Pro Cys Glu Cys Leu Val 130 Val Arg Val Ala Pro Asp Leu Gly Glu Arg Ile Thr Leu Ser Gly Asp 130 Lys Ser Leu Ile Glu Glu Val Phe Pro Glu Ile Gly Asp Val Met Cys 170 Asn Ser Val Asn Ala Gly Trp Asn His Asp Ser Thr His Val Ile Arg 180 Phe Pro Leu Asn Gly Tyr Cys His Leu Asn Ser Val Gln Val Leu Glu 200 205 Arg Leu Gln Gln Arg Gly Phe Glu Ile Val Gly Ser Cys Gly Gly Gly 210 2215 Val Asp Ser Ser Gln Phe Ser Glu Tyr Val Leu Arg Arg Glu Leu Arg 2225 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245 220 2212 DNA 2213 H. sapiens 220 2215 C200 2215 C210 225 Arg Thr Ser Leu Ser Gly Asp 226 2210 DS 2227 C210 DS 2229 (165)(756) 2223 K+Hnov2 240 176 Ceg cag ggc gagagtt gagtcagcd agattgcacc actgcactcc agcctgggcg ggcaacccgggc ggcgaaggtt gagtgagccg agattgcacc actgcactcc actgcactcc acadgaccaccacacacacacacacacacacacacacaca	His Ile Asp Val Gly His Met Tyr Thr Ser Ser Leu Ala Thr Leu 45	
50 Fro Ile Val Leu Asp Ser Leu Lys Gln His Tyr Phe Ile Asp Arg Asp 70 619 Gln Met Phe Arg Tyr Ile Leu Asn Phe Leu Arg Thr Ser Lys Leu 90 85 60 90 86 Leu Ile Pro Asp Asp Phe Lys Asp Tyr Thr Leu Leu Tyr Glu Glu Ala 100 Lys Tyr Phe Gln Leu Gln Pro Met Leu Leu Glu Met Glu Arg Trp Lys 125 115 120 Gln Asp Arg Glu Thr Gly Arg Phe Ser Arg Pro Cys Glu Cys Leu Val 130 Val Arg Val Ala Pro Asp Leu Gly Glu Arg Ile Thr Leu Ser Gly Asp 150 Lys Ser Leu Ile Glu Glu Val Phe Pro Glu Ile Gly Asp Val Met Cys 165 Lys Ser Leu Ile Glu Glu Val Phe Pro Glu Ile Gly Asp Val Met Cys 165 Asn Ser Val Asn Gly Trp Asn His Asp Ser Thr His Val Ile Arg 190 Phe Pro Leu Asn Gly Tyr Cys His Leu Asn Ser Val Gln Val Leu Glu 205 Arg Leu Gln Gln Arg Gly Phe Glu Ile Val Gly Ser Cys Gly Gly Gly 215 Val Asp Ser Ser Gln Phe Ser Glu Tyr Val Leu Arg Arg Glu Leu Arg 220 230 230 231 Val Asp Ser Ser Gln Phe Ser Glu Tyr Val Leu Arg Arg Glu Leu Arg 245 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245 220 221> CDS 221> CDS 222> (165)(756) 2223> K+Hnov2 240> 15 gogtggtggc aggtgcctgt aggcccagct acttgggagg ctgaggagg agaatagctt gagaccoggag acctcatctc aaaaaaaaag gtagttatgg ccac actg acctgaggcagagacgagac	Thr Lys Tyr Pro Glu Ser Arg Ile Gly Arg Leu Phe Asp Gly Thr Glu	
Gly Gln Met Phe Arg Tyr Ile Leu Asn Phe Leu Arg Thr Ser Lys Leu 95 85 Leu Ile Pro Asp Asp Phe Lys Asp Tyr Thr Leu Leu Tyr Glu Glu Ala 100 100 105 115 115 116 117 118 119 119 119 119 110 110 110 110 110 110	50 55 Bro He Val Leu Asp Ser Leu Lys Gln His Tyr Phe He Asp Arg Asp 80	
Leu Ile Pro Asp Asp Phe Lys Asp Tyr Thr Leu Leu Tyr Glu Glu Ala 100 105 125 120 125 125 126 135 140 135 140 135 140 135 160 155 160 150 150 150 150 150 150 150 150 150 15	65 70 75 Leu Asn Phe Leu Arg Thr Ser Lys Leu	
Lys Tyr Phe Gln Leu Gln Pro Met Leu Leu Glu Met Glu Arg Trp Lys 115 120 121 121 121 121 121 122 Gln Asp Arg Glu Thr Gly Arg Phe Ser Arg Pro Cys Glu Cys Leu Val 135 136 137 140 138 Val Arg Val Ala Pro Asp Leu Gly Glu Arg Ile Thr Leu Ser Gly Asp 145 145 155 160 145 145 150 148 155 160 155 160 155 160 175 165 170 175 180 Phe Pro Leu Asn Ala Gly Trp Asn His Asp Ser Thr His Val Ile Arg 180 180 Phe Pro Leu Asn Gly Tyr Cys His Leu Asn Ser Val Gln Val Leu Glu 205 195 195 180 Phe Glu Ile Val Gly Ser Cys Gly Gly Gly 210 211 220 Val Asp Ser Ser Gln Phe Ser Glu Tyr Val Leu Arg Arg Arg Glu Leu Arg 225 230 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245 221 222 223 234 237 240 221 238 240 221 239 241 255 221 240 221 255 221 257 261 261 271 272 284 273 275 275 276 271 276 271 277 278 278 279 270 270 271 270 270 271 270 270 271 271 272 272 273 274 275 277 277 278 278 278 278 278 278 278 278	90 95 85 90 Tyr Thr Leu Leu Tyr Glu Glu Ala	
Gln Asp Arg Glu Thr Gly Arg Phe Ser Arg Pro Cys Glu Cys Leu Val 130 Val Arg Val Ala Pro Asp Leu Gly Glu Arg Ile Thr Leu Ser Gly Asp 150 Lys Ser Leu Ile Glu Glu Val Phe Pro Glu Ile Gly Asp Val Met Cys 165 Asn Ser Val Asn Ala Gly Trp Asn His Asp Ser Thr His Val Ile Arg 180 Phe Pro Leu Asn Gly Tyr Cys His Leu Asn Ser Val Gln Val Leu Glu 205 Arg Leu Gln Gln Arg Gly Phe Glu Ile Val Gly Ser Cys Gly Gly Gly 210 Val Asp Ser Ser Gln Phe Ser Glu Tyr Val Leu Arg Arg Glu Leu Arg 225 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245 220 2210 2210 225 Arg Thr Ser Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245 220 2210 2210 2210 2210 230 230	Leu Ile Pro Asp Asp Pne Lys Asp 172 110 110 105 10 100 105 110 Met Glu Arg Trp Lys	
Val Arg Val Ala Pro Asp Leu Gly Glu Arg Ile Thr Leu Ser Gly Asp 150 150 150 150 150 150 150 150 170 175 175 175 175 165 170 175 175 175 175 175 175 175 175 175 175	Lys Tyr Phe Gln Leu Gln Pro Met Leu Glu 125 120 125 127 128 129 129 120 125 120 125 127 128 129 129 120 120 120 120 120 120	
Val Arg Val Ala Pro Asp Leu Gly Glu Arg fle Thr Leu Ser Gly Asp 150 145 Lys Ser Leu Ile Glu Glu Val Phe Pro Glu Ile Gly Asp Val Met Cys 165 Asn Ser Val Asn Ala Gly Trp Asn His Asp Ser Thr His Val Ile Arg 180 Phe Pro Leu Asn Gly Tyr Cys His Leu Asn Ser Val Gln Val Leu Glu 205 Arg Leu Gln Gln Arg Gly Phe Glu Ile Val Gly Ser Cys Gly Gly Gly 210 Val Asp Ser Ser Gln Phe Ser Glu Tyr Val Leu Arg Arg Glu Leu Arg 240 225 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245	Gln Asp Arg Glu Thr Gly Arg Phe Ser Arg Pro Cys Glu C/S 200 140	
Lys Ser Leu Ile Glu Glu Val Phe Pro Glu Ile Gly Asp Val Met Cys 165 Asn Ser Val Asn Ala Gly Trp Asn His Asp Ser Thr His Val Ile Arg 180 Phe Pro Leu Asn Gly Tyr Cys His Leu Asn Ser Val Gln Val Leu Glu 200 Arg Leu Gln Gln Arg Gly Phe Glu Ile Val Gly Ser Cys Gly Gly Gly 210 Val Asp Ser Ser Gln Phe Ser Glu Tyr Val Leu Arg Arg Glu Leu Arg 225 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245 220 2215 220 2216 2217 2218 2220 2231 Arg Thr Ser Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245 225 226 227 227 228 229 2212 DNA 2213 H. sapiens 220 2213 CDS 2222 (165)(756) 2223 K+Hnov2 240 255 260 275 286 287 288 298 298 298 298 298 298	Val Arg Val Ala Pro Asp Leu Gly Glu Arg Ile Thr Leu Ser Gly Asp 155 160	
Asn Ser Val Asn Ala Gly Trp Asn His Asp Ser Thr His Val Ile Arg 180 185 190 Phe Pro Leu Asn Gly Tyr Cys His Leu Asn Ser Val Gln Val Leu Glu 205 Arg Leu Gln Gln Arg Gly Phe Glu Ile Val Gly Ser Cys Gly Gly Gly 210 210 211 Val Asp Ser Ser Gln Phe Ser Glu Tyr Val Leu Arg Arg Glu Leu Arg 225 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245 220 221> 221> 220 221> 221> 220 221> 220 221> 220 221> 220 221> 220 221> 220 221> CDS 222> (165)(756) 2223> K+Hnov2 240 15 gegtggtgge aggtgectgt agecccaget acttgggagg ctgaggcagg agaatagett gaacccgggc ggcgaaggtt gagtgagceg agattgcacc actgcactcc agectgggeg gaacccgggc ggcgaaggtt gagtgagceg agattgcacc actgcactcc agectgggeg gaacccgggc ggcgaaggtt gagtgagceg agattgcacc actgcactcc agectgggeg acagagcgag actccatctc aaaaaaaaaga gtagttatgg ccac atg gcc cca cta Met Ala Pro Leu 1 tcg cca ggc gga aag gcc ttc tgc atg gtc tat gca gcc ctg ggg ctg Ser Pro Gly Gly Lys Ala Phe Cys Met Val Tyr Ala Ala Leu Gly Leu 15	145 Lyc Ser Leu Ile Glu Glu Val Phe Pro Glu Ile Gly Asp Val Met Cys 175	
Phe Pro Leu Asn Gly Tyr Cys His Leu Asn Ser Val Gln Val Leu Glu 200 Arg Leu Gln Gln Arg Gly Phe Glu Ile Val Gly Ser Cys Gly Gly Gly 210 210 Val Asp Ser Ser Gln Phe Ser Glu Tyr Val Leu Arg Arg Glu Leu Arg 220 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245 220 235 240 255 255 255 255 255 220 255 <a 20.15"="" href="https://doi.org/10.15">255 <a 20.15"="" 2<="" href="https://doi.org/10.15" td=""><td>165 170 165 Trp Asn His Asp Ser Thr His Val Ile Arg</td><td></td>	165 170 165 Trp Asn His Asp Ser Thr His Val Ile Arg	
Arg Leu Gln Gln Arg Gly Phe Glu Ile Val Gly Ser Cys Gly Gly Gly 210 215 Val Asp Ser Ser Gln Phe 230 231 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245 245 250 2610 275 286 287 287 288 298 298 298 298 298	Asn Ser val Ash Alta Say 185 185 180 180 180 180 180 Asn Ser Val Gln Val Leu Glu	
Val Asp Ser Ser Gln Phe Ser Glu Tyr Val Leu Arg Arg Glu Leu Arg 235 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 245 246 257 240 258 240 259 250 255 260 275 275 286 287 288 298 298 298 298 298 298	Phe Pro Leu Asn Gly Tyr Cys his account 205	
Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 255 Color	Arg Leu Gln Gln Arg Gly Phe Glu 11e var Gly 321 3220	
Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 255 Color	Val Asp Ser Ser Gln Phe Ser Glu Tyr Val Leu Arg Arg Glu Leu Arg 240	
<pre></pre>	225 230 Arg Thr Pro Arg Val Pro Ser Val Ile Arg Ile Lys Gln Glu Pro Leu 255 255	
<pre> <211> 923 <212> DNA <213> H. sapiens <220> <221> CDS <222> (165)(756) <223> K+Hnov2 </pre> <pre> <</pre>	245	
<pre> <212> DNA <213> H. sapiens <220></pre>		
<pre></pre>	<212> DNA	
<pre> <221> CDS <222> (165)(756) <223> K+Hnov2 </pre> <pre> <400> 15 gcgtggtggc aggtgcetgt agccccaget acttgggagg ctgaggcagg agaatagett gagtgggcg agattgcacc actgcactcc agcctgggcg gaacccgggc ggcgaaggtt gagtgagccg agattgcacc actgcactcc agcctgggcg acacagggag actccatctc aaaaaaaaaa</pre>	<213> H. sapiens	
<pre> <222> (165)(756) <223> K+Hnov2 <400> 15 gcgtggtggc aggtgcetgt agccccagct acttgggagg ctgaggcagg agaatagctt gaacccgggc ggcgaaggtt gagtgagccg agattgcacc actgcactcc agcctgggcg acagggaggaggaggaggaggaggaggaggaggaggagga</pre>		
<223> K+Hnov2 <400> 15 gcgtggtggc aggtgcctgt agccccagct acttgggagg ctgaggcagg agaatagctt gagtggcg aggtgccgg agattgcacc actgcactcc agcctgggcg agaacccgggc ggcgaaggtt gagtgagccg agattgcacc actgcactcc agcctgggcg acacagggcgag actccatctc aaaaaaaaaga gtagttatgg ccac atg gcc cca cta Met Ala Pro Leu 1 tcg cca ggc gga aag gcc ttc tgc atg gtc tat gca gcc ctg ggg ctg Ser Pro Gly Gly Lys Ala Phe Cys Met Val Tyr Ala Ala Leu Gly Leu 224	<222> (165)(756)	
gegtggtggc aggtgcetgt agceccaget acttgggagg ctgaggcagg agatagets gaaccegggc ggcgaaggtt gagtgagceg agattgcacc actgcactcc agcetgggeg acaggggga actccatctc aaaaaaaaga gtagttatgg ccac atg gcc cca cta Met Ala Pro Leu 1 teg cca ggc gga aag gcc ttc tgc atg gtc tat gca gcc ctg ggg ctg Ser Pro Gly Gly Lys Ala Phe Cys Met Val Tyr Ala Ala Leu Gly Leu 224	<223> K+Hnov2	
gaacceggge ggegaaggtt gagtgagees system acagagegag actecatete aaaaaaaaga gtagttatgg ccac atg gee cca cta Met Ala Pro Leu 1 teg cca gge gga aag gee tte tge atg gte tat gea gee etg ggg etg Ser Pro Gly Gly Lys Ala Phe Cys Met Val Tyr Ala Ala Leu Gly Leu 224	<400> 15	
tcg cca ggc gga aag gcc ttc tgc atg gtc tat gca gcc ctg ggg ctg tcg rpro Gly Gly Lys Ala Phe Cys Met Val Tyr Ala Ala Leu Gly Leu 1 224	gegtggtggc aggtgcctgt agetecases agattgcacc actgcactcc agcctgggcg	-
tog coa ggc gga aag gcc tto tgc atg gtc tat gca gcc ctg ggg cos Ser Pro Gly Gly Lys Ala Phe Cys Met Val Tyr Ala Ala Leu Gly Leu	acagagegag actecatete adadudadys 500 Met Ala Pro Leu	
Ser Pro Gly Gly Lys Ala File Cys 15	are are are the type aty gto tat goa goe etg ggg etg	224
	Ser Pro Gly Gly Lys Ala File Cys	
5	5	272
cca gcc tcc tta gct ctc gtg gcc acc ctg cgc cat tgc ctg ctg cct 272	cca ged ted tha get che gig ged acc eng ege tad ege too 500 000 000 000 000 000 000 000 000 0	

Pro Ala Ser Leu Ala Leu Val Ala Thr Leu Arg His Cys Leu Leu Pro	
gtg ctc agc cgc cca cgt gcc tgg gta gcg gtc cac tgg cag ctg tca gtg ctc agc cgc cca cgt gcc tgg gta gcg gtc cac tgg cag ctg tca 3 Val Leu Ser Arg Pro Arg Ala Trp Val Ala Val His Trp Gln Leu Ser 45 50	320
40	368
gcc agc agc ttt gtg ctg ctg cca gcg ctg gtg ctg tgg ggc ctt cag Ala Ser Ser Phe Val Leu Leu Pro Ala Leu Val Leu Trp Gly Leu Gln 75	416
ggc gac tgc agc ctg ctg ggg gcc gtc tac ttc tgc ttc agc tcg ctc ggc gac tgc agc ctg ctg ggg gcc gtc tac ttc tgc ttc agc tcg ctc ggc gac tgc agc ctg ctg ggg gcc gtc tac ttc tgc ttc agc tcg ctc ggc gac tgc agc ctg ctg ggg gcc gtc tac ttc tgc ttc agc tcg ctc ggc gac tgc agc ctg ctg agc gtc tac ttc tgc ttc agc tcg ctc ggc gac tgc agc tcg ctc ggc gac tgc agc tcg ctc ggc gac tgc agc tgc tac ttc tgc ttc agc tcg ctc ggc gac tgc agc tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc ggc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc gtc tgc agc gtc tac ttc tgc ttc agc tcg ggc gac tgc agc gtc tgc agc gtc tac ttc tgc tcg ggc gac tgc agc gtc tgc agc gtc tgc tcg ggc gac tgc agc gtc tgc agc gtc tac ttc tgc tcg ggc gac tgc agc gtc tgc agc gtc tac ttc tgc tcg ggc gac tgc agc gtc tgc agc gtc tgc tgc tgc tgc tgc tgc tgc tgc t	464
agc acc att ggc ctg gag gac ttg ctg ccc ggc cgc ggc cgc agc ctg agc acc att ggc ctg gag gac ttg ctg ccc ggc cgc ggc cgc agc ctg agc acc att ggc ctg gag gac ttg ctg ccc ggc cgc ggc cgc agc ctg agc acc att ggc ctg gag gac ttg ctg ccc ggc cgc ggc cgc agc ctg agc acc att ggc ctg gag gac ttg ctg ccc ggc cgc ggc cgc agc ctg agc acc att ggc ctg gag gac ttg ctg ccc ggc cgc ggc cgc agc ctg	512
cac ccc gtg att tac cac ctg ggc cag ctc gca ctt ctt ggt tac ttg	560
ctt cta gga ctc ttg gcc atg ctg ctg gca gtg gag acc ttc tct gag ctt cta gga ctc ttg gcc atg ctg ctg squ gag acc ttc tct gag Leu Leu Gly Leu Leu Ala Met Leu Leu Ala Val Glu Thr Phe Ser Glu 140 145	608
ctg ccg cag gtc cgt gcc atg ggg aag ttc ttc aga ccc agt ggt cct	656
gtg act gct gag gac caa ggt ggc atc cta ggg cag gat gaa ctg gct yal Thr Ala Glu Asp Gln Gly Gly Ile Leu Gly Gln Asp Glu Leu 170 175 180	704
ctg agc acc ctg ccg ccc gcg gcc cca gct tca gga caa gcc cct gct ctg Ser Thr Leu Pro Pro Ala Ala Pro Ala Ser Gly Gln Ala Pro Ala Leu Ser Thr Leu Pro Pro Ala Ala Pro Ala Ser Gly Gln Ala Pro Ala	752
tgc t gaagcgtcag gtgaccgagt tcagctccgt aaggtggcgg cacctgagga	806
Cys ggaagcagcc aggagtggct ggggaagaat ctggagatgg agccgcggtg agggtgggcg ggaggcctca ggggatactg ttaatcataa aaaaaaaaaa	866 923
<210> 16 <211> 197 <212> PRT <213> H. sapiens	
<pre> <400> 16 Met Ala Pro Leu Ser Pro Gly Gly Lys Ala Phe Cys Met Val Tyr Ala</pre>	
26	

40 45	
35 Trp Gln Leu Ser Pro Ala Arg Ala Ala Leu Leu Gln Ala Val Ala Leu 55 60 77 78 78 78 78 78 78 78 78 78 78 78 78	
55 50 Solvation	
Gly Leu Leu Val Ala Sel Sel 110 75 75 70 70 70 70 71 71 71 75 75 76 75 76 77 76 77 77 78 78 78 78 78 78 78 78 78 78 78	
65 70 75 65 Trp Gly Leu Gln Gly Asp Cys Ser Leu Leu Gly Ala Val Tyr Phe Cys 95 95	
90 85 85 Phe Ser Ser Leu Ser Thr Ile Gly Leu Glu Asp Leu Leu Pro Gly Arg 110 105 105 107 108 109 110	
105 100 Gly Arg Ser Leu His Pro Val Ile Tyr His Leu Gly Gln Leu Ala Leu 125 120 120 125 120 125 120 127 128 129 120 120 121 125	
120 115 120 Leu Gly Tyr Leu Leu Leu Gly Leu Leu Ala Met Leu Leu Ala Val Glu 140 140 140 140 140 140 140 140 140 140	
Leu Gly Tyr Leu Leu Leu Gly Heu 135 130 130 130 140 135 140 130 130 140 130 130 130 130	
135 130 135 136 137 138 139 130 130 135 140 140 140 140 140 140 140 140 140 140	
Thr Phe Set State 150 155 156 157 158 159 159 150 150 155 170 175 170 175 170 175 175 170 175 175 170 175 175 175 175 175 175 175 175 175 175	
Pro Ser Gly Pro Val III 120 170 170 170 Pro Ala Ser Gly	
Asp Glu Leu Ala Leu Ser Thr Leu Pro Pro Ala Ala Pro Ala Ser Gly 185 185	
180 Gln Ala Pro Ala Cys	
195	
<210> 17	
<211> 3102	
<212> DNA <213> H. sapiens	
<213> π. σωμ2σαστ	
<220>	
<221> CDS <222> (274)(1705)	
<223> K+Hnov11	
2223 RTIMO 12-	
A005 17	60
<400> 17	120
<400> 17 gcacgegcaa agegeceaec gagacceetg gggtggaget tgtgttaata gaaacatace gcacgegcaa agegeceaec gagacceetg agecetcaaa etettgecec ageceageec	120 180
<400> 17 gcacgegcaa agegcccacc gagacccctg gggtggagct tgtgttaata gaaacatacc gcacgegcaa cttcctggg aggggatcag acccctcaaa ctcttgccc agcccagccc	120
<400> 17 gcacgegcaa agegcccacc gagacccctg gggtggagct tgtgttaata gaaacatacc gcacgegcaa cttcctggg aggggatcag acccctcaaa ctcttgccc agcccagccc	120 180 240
<400> 17 gcacgegcaa agegeceaec gagacceetg gggtggaget tgtgttaata gaaacatace gcacgegcaa agegeceaec gagacceetg agecetcaaa etettgecec ageceageec	120 180 240
caccecage ctttcctggg aggagactetg gggtggaget tgtgttaata gaaacatace caccecage ctttcctggg aggggatcag accettaaa ctettgecee agecaggegggggggggggggggggggggggggg	120 180 240
caccecage ctttcctggg aggagactetg gggtggaget tgtgttaata gaaacatace caccecage ctttcctggg aggggatcag accettaaa ctettgecee agecaggegggggggggggggggggggggggggg	120 180 240 294
geacgegeaa agegeeeace gagaceetg gggtggaget tgtgttaata gaaacatace cacececage ettteetggg aggggateag acceeteaaa etettgeeee ageeaggggtteageee cagaceece aagaceeace aggaggeetg ggeeegeeag taatgggtag ggagaggggg etetegeeag egetgtteee teegetteea ggtgtagege egeegeege egeggeetee age atg ace gge cag age etg tgg egegeetee age atg ace gge cag age etg tgg het Thr Gly Gln Ser Leu Trp 1 5 gac gtg tcg gag get aac gte gag gac ggg gag ate ege ate aat gtg Asp Val Ser Glu Ala Asn Val Glu Asp Gly Glu Ile Arg Ile Asn Val	120 180 240 294
geacgegeaa agegeceaec gagacecetg gggtggaget tgtgttaata gaaacatace caceceage ettreetggg agggateag acceeteaaa etettgeece agecageee tteageaece aagaceaeca aggaggeetg ggceegeag taatgggtag ggagaggggg etetegeega egetgttee teegetteea ggtgtagege eeceegeegg egeggeegee eggegeetee age at acc gge eag age etg tgg eeceegeegg egeggeetee eggegeetee age at acc gge eag age etg tgg age ggg gag at ee ege at eat gtg gac gtg teg gag get aac gte gag gac ggg gag at eege at aat gtg Asp Val Ser Glu Ala Asn Val Glu Asp Gly Glu Ile Arg Ile Asn Val 10 15	120 180 240 294
geacgegeaa agegeceaec gagacecetg gggtggaget tgtgttaata gaaacatace caceceage ettreetggg agggateag acceeteaaa etettgeece agecageee tteageaece aagaceaeca aggaggeetg ggceegeag taatgggtag ggagaggggg etetegeega egetgttee teegetteea ggtgtagege eeceegeegg egeggeegee eggegeetee age at acc gge eag age etg tgg eeceegeegg egeggeetee eggegeetee age at acc gge eag age etg tgg age ggg gag at ee ege at eat gtg gac gtg teg gag get aac gte gag gac ggg gag at eege at aat gtg Asp Val Ser Glu Ala Asn Val Glu Asp Gly Glu Ile Arg Ile Asn Val 10 15	120 180 240 294
geacgegeaa agegeeeace gagacceetg gggtggaget tgtgttaata gaaacatace cacceceage ettteetggg agggggateag acceeteaaa etettgeeee ageagaggegg tteageacee aagaceeace aggaggeetg ggeeegeeag taatgggtag gggagaggggg etetegeeag egetgtteee teegetteea ggtgtagege egegegeetee age atg ace gge cag age etg tgg ceegegegg egegegeetee eggegeetee age atg ace gge cag age etg tgg Gge gge ggggeetee age atg ace gge cag age etg tgg Gge ggg gag ate ege ate aat gtg as par se ggg gag ate ege ate aat gtg as par se ggg gag ate ege ate aat gtg as par se ggg ggg gag ate ege ate aat gtg as par se ggg ggg gag ate ege ate aat gtg as par se ggg ggg ggg ggg ette aat gtg ag ggg ggg ggg ate ege ate aat gtg as par se ggg ggg ggg ggg ate ege ate aat gtg ag ggg ggg ggg ggg ate ege ate aat gtg ag ggg ggg ggg ggg ate ege ate aat gtg ag ggg ggg ggg ggg ate ege ate aat gtg ag ggg ggg ggg ggg ag ate ege ate aat gtg ag ggg ggg ggg ggg ag ate ege ate aat gtg ag ggg ggg ggg ggg ag ate ege ate aat gtg ag ggg ggg ggg ggg ggg ag ate ege ate aat gtg ag ggg ggg ggg ggg ggg ggg ggg	120 180 240 294 342
geacgegeaa agegeeeace gagacceetg gggtggaget tgtgttaata gaaacatace cacceceage ettteetggg agggggateag acceeteaaa etettgeeee ageeagggggtteageee eageegeeag ggeeegeeag taatgggtag gggagaggggg eteteegeeag egetgtteee teegetteea ggtgtagege egeegeegeg egegegeetee age atg ace gge eag age etg tgg ceegeegegg egegegeetee eggegeetee age atg ace gge eag age etg tgg egeegeegegg eggegeetee eggegeetee age atg ace gge eag age etg tgg egge eggegeetee age atg ace gge eag age etg tgg fall fall fall fall fall fall fall fa	120 180 240 294
geacgegeaa agegeeeace gagacceetg gggtggaget tgtgttaata gaaacatace cacceceage ettteetggg agggggateag acceeteaaa etettgeeee ageeagggggtteageee eageegeeag ggeeegeeag taatgggtag gggagaggggg eteteegeeag egetgtteee teegetteea ggtgtagege egeegeegeg egegegeetee age atg ace gge eag age etg tgg ceegeegegg egegegeetee eggegeetee age atg ace gge eag age etg tgg egeegeegegg eggegeetee eggegeetee age atg ace gge eag age etg tgg egge eggegeetee age atg ace gge eag age etg tgg fall fall fall fall fall fall fall fa	120 180 240 294 342
geacgegeaa agegeeeace gagaceetg gggtggaget tgtgttaata gaaacatace cacceceage etttectggg aggggateag accecteaaa etettgeeee ageeageeg teageacee aagaceeace aggaggeetg ggceegeag taatgggtag ggagaggggg etetegeag egetgtteee teegetteea ggtgtaageg eeeegeegg egegegege eggegeetee age atg ace gge cag age etg tgg ceegeegg eggegeetee age atg ace gge cag age etg tgg eggegegege eggegeetee age atg ace gge cag age etg tgg eggegegegegegegegegegegegegeg	120 180 240 294 342 390
geacgegeaa agegeeeace gagaceetg gggtggaget tgtgttaata gaaacatace cacececage etttectggg aggggateag accecteaaa etettgeee ageeaggeg teageaceage egetgtteee teegetteea ggggagagggggeeegeegeegegegegegegegeg	120 180 240 294 342
geacgegeaa agegeeeace gagaceetg gggtggaget tgtgttaata gaaacatace cacececage etttectggg aggggateag accecteaaa etettgeee ageeaggeg teageaceage egetgtteee teegetteea ggggagagggggeeegeegeegegegegegegegeg	120 180 240 294 342 390
geacgegeaa agegeceaec gagacecetg gggtggaget tgtgttaata gaaacatace cacececage etteetggg agggagetag acceeteaaa etettgeee agecageee teageaece aagaceaece aggaggeetg ggceegeeag taatgggtag ggagaggggg eteteegeea eggetgtteee teegetteea ggtgtagege eegeegeegege eegeegeeege eggegeetee eggegeetee age at aatgggtag ggagaggggg eegeegeegeegeegeegeegeegeegeegeege	120 180 240 294 342 390 438
geacgcgaaa agcgccaacc gagacccctg gggtggaget tgtgttaata gaaacatacc caccccage ctttectggg aggggatcag accectcaaa ctcttgcccc agccagccc caccccagc ctttectggg aggaggctg ggcccgcag taatgggtag ggagaggggg cccccgccagg cgcgaccgcc ccccgccagg cgcgcccccccc	120 180 240 294 342 390
geacgegeaa agegeeeace gagacecetg gggtggaget tgtgttaata gaaacatace cacceceage ettteetggg agggagetag accectcaaa etettgeeee ageagagegegetee eegetgtteee teegetteea gggagagggggeeegeeegeegegegegegegegeg	120 180 240 294 342 390 438

ggc aag ctt cac gtc atg gct gag cta tgt gtc ttc tcc ttc agc cag Gly Lys Leu His Val Met Ala Glu Leu Cys Val Phe Ser Phe Ser Gln	582
gag atc gag tac tgg ggc atc aac gag ttc ttc att gac tcc tgc tgc Glu Ile Glu Tyr Trp Gly Ile Asn Glu Phe Phe Ile Asp Ser Cys Cys	630
agc tac agc tac cat ggc cgc aaa gta gag ccc gag cag gag aag tgg Ser Tyr Ser Tyr His Gly Arg Lys Val Glu Pro Glu Glu Lys Trp 125 130 135	678
gac gag cag agt gac cag gag agc acc acg tct tcc ttc gat gag atc gac gag cag agt gac cag gag agc acc acg tct tcc ttc gat gag atc Asp Glu Gln Ser Asp Gln Glu Ser Thr Thr Ser Ser Phe Asp Glu Ile 140 145	726
ctt gcc ttc tac aac gac gcc tcc aag ttc gat ggg cag ccc ctc ggc Leu Ala Phe Tyr Asn Asp Ala Ser Lys Phe Asp Gly Gln Pro Leu Gly 155 160 165	774
aac ttc cgc agg cag ctg tgg ctg gcg ctg gac aac ccc ggc tac tca Asn Phe Arg Arg Gln Leu Trp Leu Ala Leu Asp Asn Pro Gly Tyr Ser	822
gtg ctg agc agg gtc ttc agc atc ctg tcc atc ctg gtg gtg atg ggg Val Leu Ser Arg Val Phe Ser Ile Leu Ser Ile Leu Val Val Met Gly 190 195	870
tcc atc atc acc atg tgc ctc aat agc ctg ccc gat ttc caa atc cct tcc atc atc acc atg tgc ctc aat agc ctg ccc gat ttc caa atc cct Ser Ile Ile Thr Met Cys Leu Asn Ser Leu Pro Asp Phe Gln Ile Pro 205 210 215	918
gac agc cag ggc aac cct ggc gag gac cct agg ttc gaa atc gtg gag gac agc cag ggc aac cct ggc gag gac cct agg ttc gaa atc gtg gag Asp Ser Gln Gly Asn Pro Gly Glu Asp Pro Arg Phe Glu Ile Val Glu Asp Ser Gln Gly Asn Pro Gly Glu Asp 225	966
cac ttt ggc att gcc tgg ttc aca ttt gag ctg gtg gcc agg ttt gct His Phe Gly Ile Ala Trp Phe Thr Phe Glu Leu Val Ala Arg Phe Ala 235 240 245	1014
gtg gcc cct gac ttc ctc aag ttc ttc aag aat gcc cta aac ctt att Val Ala Pro Asp Phe Leu Lys Phe Phe Lys Asn Ala Leu Asn Leu Ile 250 255 260	1062
gac ctc atg tcc atc gtc ccc ttt tac atc act ctg gtg gtg aac ctg Asp Leu Met Ser Ile Val Pro Phe Tyr Ile Thr Leu Val Val Asn Leu 270 275	1110
gtg gtg gag agc aca cct act tta gcc aac ttg ggc agg gtg gcc cag Val Val Glu Ser Thr Pro Thr Leu Ala Asn Leu Gly Arg Val Ala Gln Val Val Glu Ser Thr Pro Thr Leu Ala Asn Leu Gly Arg Val Ala Gln 295	1158
gtc ctg agg ctg atg cgg atc ttc cgc atc tta aag ctg gcc agg cac gtc ctg agg ctg atg cgg atc ttc cgc atc tta aag ctg gcc agg cac Val Leu Arg Leu Met Arg Ile Phe Arg Ile Leu Lys Leu Ala Arg His 300 305 310	1206
tcc act ggc ctc cgc tcc ctg ggg gcc act ttg aaa tac agc tac aaa Ser Thr Gly Leu Arg Ser Leu Gly Ala Thr Leu Lys Tyr Ser Tyr Lys	1254
gaa gta ggg ctg ctc ttg ctc tac ctc tcc gtg ggg att tcc atc ttc	1302

Glu Val Gly Leu Leu Leu Tyr Leu Ser Val Gly Ile Ser Ile Phe	
330 and gag gag gag ctq gcc 13	50
tcc gtg gtg gcc tac acc att gaa aag gag gag aac gag gag Ser Val Val Ala Tyr Thr Ile Glu Lys Glu Glu Asn Glu Gly Leu Ala 345 350 355	,
	398
ggg tac ggg gat gtg gtc cca ggg acc acg gca gga aag ctg act gcc 1. Gly Tyr Gly Asp Val Val Pro Gly Thr Thr Ala Gly Lys Leu Thr Ala 380 385 390	446
tct gcc tgc atc ttg gca ggc atc ctc gtg gtg gtc ctg ccc atc acc 1 Ser Ala Cys Ile Leu Ala Gly Ile Leu Val Val Leu Pro Ile Thr 395 400 405	494
	L542
gag agt gcc atg cgc agc tgt gac ttt gga gat gga atg aag gag gtc Glu Ser Ala Met Arg Ser Cys Asp Phe Gly Asp Gly Met Lys Glu Val 425 430 435	1590
cct tcg gtc aat tta agg gac tat tat gcc cat aaa gtt aaa tcc ctt Pro Ser Val Asn Leu Arg Asp Tyr Tyr Ala His Lys Val Lys Ser Leu 455	1638
atg gca agc ctg acg aac atg agc agg agc tca cca agt gaa ctc agt atg gca agc ctg acg aac atg agc agg agc tca cca agt gaa ctc agt atg gca agc ctg acg acg agc ser Pro Ser Glu Leu Ser Met Ala Ser Leu Thr Asn Met Ser Arg Ser Pro Ser Glu Leu Ser 460 460	1686
tta aat gat tcc cta cgt t agecgggagg acttgtcacc ctccacccca Leu Asn Asp Ser Leu Arg 475	1735
	1795
cattgctgag ctgcctcttg tgcctctggc acagcccagg caccttatgg ttatggtgta	1855
aggagtatge ccagecetty aggggagaga and an aggregatett geacettece	1915 1975
cagtttttag aatcgttttt agaggggss s s s stattcacct tttttccaga	2035
atgaaatgac actcactggt ctttgcatcg tgggcataaa atgtcttcta gtgcttgtgg tgagtacacc cagaatgcta attttctgt ccatcgtgta cgctattcta gtgcttgtgt tgagtacacc cagaatgcta attttctgt tgttctqag gttgtcgtgt gagttctgta	2095
annatacta totatqagtt gicgigotto si annatagga aggatatqat	2155
caaaaagccc ccacaagtcg tttagtcgut ctttatccat tqctttcact	2215
gagattttgc tcacagtcat gryanactur throngs gagttgacca atgcaagtct	2275 2335
tagttttagt accadaded against a categgitgea ccaatcacce	2395
ctaagttgtt tttataaatg atctgtagtt ccgtggtttg catssssssssssssssssssssssssssssssssssss	2455
gttaaacatg cagtgaagat cgautettaatga agttagtgca ggctgagccc	2515
cttcagctaa atagtccada ccctaggg	2575 2635
cttogttcac agtcaageet coossississississississississississississ	2695
atgaggtttc tgatctaggc catttgacca aactttgctg tgetadstatecta ttttgaaata tttattttt aagatgttta ggagtaaaggt cgtgttgtct tcctcaacta ttttgaaata tttattttt aagatgttta ggagtgaacg ttgttgggtt gctagcaagg	2755
agaagaagtt tactgttgta tegetteet sangagage cettttattt	2815
cartagetta atactitigi tyeettasta agratattat atatgatget	2875 2935
ccaagcagaa tttagtcaga taatttagaa tagtatagaa qcatgtactc	2935 2995
gtgattgccc tggagttcct gcccatgact ggaaacctgg tggaaacac tgcagttctt aaaatataga cgtgcacgat ggtggtgtgg cttacccagg atggaaacac tgcagttctt acttgcattc ccactgcctt tcatgggggg tgactgggta gaggccagga gaaaggaaag	3055
pottagatta ccactacett ccacdadada carridge	

3102

<210> 18 <211> 477 <212> PRT

<213> H. sapiens

Met Thr Gly Gln Ser Leu Trp Asp Val Ser Glu Ala Asn Val Glu Asp Gly Glu Ile Arg Ile Asn Val Gly Gly Phe Lys Arg Arg Leu Arg Ser 25 His Thr Leu Leu Arg Phe Pro Glu Thr Arg Leu Gly Arg Leu Leu Leu Cys His Ser Arg Glu Ala Ile Leu Glu Leu Cys Asp Asp Tyr Asp Asp 40 Val Gln Arg Glu Phe Tyr Phe Asp Arg Asn Pro Glu Leu Phe Pro Tyr

Val Leu His Phe Tyr His Thr Gly Lys Leu His Val Met Ala Glu Leu

Cys Val Phe Ser Phe Ser Gln Glu Ile Glu Tyr Trp Gly Ile Asn Glu 105 Phe Phe Ile Asp Ser Cys Cys Ser Tyr Ser Tyr His Gly Arg Lys Val

Glu Pro Glu Gln Glu Lys Trp Asp Glu Gln Ser Asp Gln Glu Ser Thr 130 135 140

Thr Ser Ser Phe Asp Glu Ile Leu Ala Phe Tyr Asn Asp Ala Ser Lys
155
160

Phe Asp Gly Gln Pro Leu Gly Asn Phe Arg Arg Gln Leu Trp Leu Ala 175 Leu Asp Asn Pro Gly Tyr Ser Val Leu Ser Arg Val Phe Ser Ile Leu

Ser Ile Leu Val Val Met Gly Ser Ile Ile Thr Met Cys Leu Asn Ser

Leu Pro Asp Phe Gln Ile Pro Asp Ser Gln Gly Asn Pro Gly Glu Asp

Pro Arg Phe Glu Ile Val Glu His Phe Gly Ile Ala Trp Phe Thr Phe

Glu Leu Val Ala Arg Phe Ala Val Ala Pro Asp Phe Leu Lys Phe Phe 245 250

Lys Asn Ala Leu Asn Leu Ile Asp Leu Met Ser Ile Val Pro Phe Tyr

Ile Thr Leu Val Val Asn Leu Val Val Glu Ser Thr Pro Thr Leu Ala 280

Asn Leu Gly Arg Val Ala Gln Val Leu Arg Leu Met Arg Ile Phe Arg 295

Ile Leu Lys Leu Ala Arg His Ser Thr Gly Leu Arg Ser Leu Gly Ala 305 310 315 Thr Leu Lys Tyr Ser Tyr Lys Glu Val Gly Leu Leu Leu Tyr Leu

Ser Val Gly Ile Ser Ile Phe Ser Val Val Ala Tyr Thr Ile Glu Lys

345 Glu Glu Asn Glu Gly Leu Ala Thr Ile Pro Ala Cys Trp Trp Trp Ala 360

Thr Val Ser Met Thr Thr Val Gly Tyr Gly Asp Val Val Pro Gly Thr

Thr Ala Gly Lys Leu Thr Ala Ser Ala Cys Ile Leu Ala Gly Ile Leu

Val Val Val Leu Pro Ile Thr Leu Ile Phe Asn Lys Phe Ser His Phe

Tyr Arg Arg Gln Lys Gln Leu Glu Ser Ala Met Arg Ser Cys Asp Phe 30

425	
420 Gly Asp Gly Met Lys Glu Val Pro Ser Val Asn Leu Arg Asp Tyr Tyr 445 440 440 445 440 440 445 447	
Gly Asp Gly Met Lys Gat 440 435 Ala His Lys Val Lys Ser Leu Met Ala Ser Leu Thr Asn Met Ser Arg 460 455	
Ala His Lys Val Lys 455 455 450 Asn Asp Ser Leu Arg	
455 450 Ser Ser Pro Ser Glu Leu Ser Leu Asn Asp Ser Leu Arg 475 465	
<210> 19 <211> 0 <212> DNA <213> H. sapiens	
<220> <221> CDS <222> (249)(3495) <223> K+Hnovl4	60
<400> 19 gggctggtag cagggatttg tgggcggcga gggcgcgagg ggccgcgcgc catgctccgg gggctggtag cagggatttg tgggcggcg ccagctccgg gcgaccccg gatcccggtc gggctggtag caggaggc ccctcgcgcg ccagctccgg gcgacgagct	60 120
gggctggtag cagggatttg tgggcggcga gggcgcgagg ggccgcggg gatcccggtc gatcccggtc gccccgacgg cgcggacgcc cctcgcgcg ccagctccgg ccggcggggg cggccgagtt gccccgacgacgc ctgcgctagg agccgcgagg cagccgcgagg	180 240
gggggccctc ccccggggg gagtccccgc accccggagg acgsssssssssssssssssss	290
1 and the gtg	338
ctg gac acc atc gct acg cgc ttc gac ggc acg cac agt aac ttc gtg ctg gac acc atc gct acg cgc ttc gac ggc acg cac agt aac ttc gtg ctg gac acc atc gct acg cgc ttc gac ggc acg cac agt aac ttc gtg ctg gac acc atc gct acg cgc ttc gac ggc acg cac agt aac ttc gtg ctg gac acc atc gtc acg cgc ttc gac ggc acg cac agt aac ttc gtg ctg gac acc atc gtc acg cac agt aac ttc gtg ctg gac acg cac acg cac agt aac ttc gtg ctg gac acg cac acg cac agt aac ttc gtg ctg gac acg cac acg cac acg cac agt aac ttc gtg ctg gac acg cac acg cac acg cac acg cac acg cac agt acg cac acg c	
15	386
ctg ggc aac gcc agt ggc ggg gct ctt ccc gtg gtc tac tgc tct gat ctg ggc aac gcc agt ggc ggg gct ctt ccc gtg gtc tac tgc tct gat ctg ggc aac gcc agt ggc ggg gct ctt ccc gtg gtc tac tgc tct gat ctg ggc aac gcc agt ggc ggg gct ctt ccc gtg gtc tac tgc tct gat ctg ggc aac gcc agt ggc ggg gct ctt ccc gtg gtc tac tgc tct gat at a ctg gcc as ac gcc agt ggc ac tac tgc tct gat ctg ggc aac gcc agt ggc ggt ctt ccc gtg gtc tac tgc tct gat at a ctg gcc as ac gcc agt ggc ggc ac ctt ccc gtg gtc tac tgc tct gat ctg ggc aac gcc agt ggc ggg gct ctt ccc gtg gtc tac tgc tct gat ac ac gcc agt ggc ggg gct ctt ccc gtg gtc tac tgc tct gat ac ac a	
ggc ttc tgt gac ctc acg ggc ttc tcc cgg gct gag gtc atg cag cgg Gly Phe Cys Asp Leu Thr Gly Phe Ser Arg Ala Glu Val Met Gln Arg	434
ggc tgt gcc tgc tcc ttc ctt tat ggg cca gac acc agt gag ctc gtc ggc tgt gcc tgc tcc ttc ctt tat ggg cca gac acc agt gag ctc gtc Gly Cys Ala Cys Ser Phe Leu Tyr Gly Pro Asp Thr Ser Glu Leu Val 70 75	482
Gly Cys Ala Cys 552 70	530
Arg Gln Gin He Als 85	570
gag ctg atc ctg tac cgg aag agc ggg ctc ccg ttc tgg tgt ctc ctg Glu Leu Ile Leu Tyr Arg Lys Ser Gly Leu Pro Phe Trp Cys Leu 100 105	578
gat gtg ata ccc ata aag aat gag aaa ggg gag gtg gct ctc ttc cta gat gtg ata ccc ata aag aat gag aaa ggg gag gtg gct ctc ttc cta Asp Val Ile Pro Ile Lys Asn Glu Lys Gly Glu Val Ala Leu Phe Leu 125	626
Asp Val Tie Fio 125 120 120 115	674
gtc tct cac aag gac atc agc gaa acc aag aac cga ggg ggc ccc gac Val Ser His Lys Asp Ile Ser Glu Thr Lys Asn Arg Gly Gly Pro Asp 135 140	
aga tgg aaa gag aca ggt ggt ggc cgg cgc cga tat ggc cgg gca cga aga tgg aaa gag aca ggt ggt ggc cgg cgc cga tat ggc cgg gca cga Arg tgg aaa gag aca ggt ggt ggc cgg cgc cga tat ggc cgg gca cga aga tgg aaa gag aca ggt ggt ggc cgg cgc cga tat ggc cgg gca cga aga tgg aaa gag aca ggt ggt ggc cgg cgc cga tat ggc cgg gca cga aga tgg aaa gag aca ggt ggt ggc cgg cgc cga tat ggc cgg gca cga aga tgg aaa gag aca ggt ggt ggc cgg cgc cga tat ggc cgg gca cga aga tgg aaa gag aca ggt ggt ggc cgg cgc cga tat ggc cgg gca cga aga tgg aaa gag aca ggt ggt ggc cgg cgc cga tat ggc cgg gca cga aga tgg aaa gag aca ggt ggt ggc cgg cgc cga tat ggc cgg gca cga aga tgg aaa gag aca ggt ggt ggc aga cga aga aga aga aga aga aga aga	722
Arg Trp Lys Glu Thr Gly 31	

155	
150	770
cac ctg tcc ggg cac ctg cag aag cac caag ggc aag cac aag ctc His Leu Ser Gly His Leu Gln Lys Gln Pro Lys Gly Lys His Lys Leu His Leu Ser Gly His Leu Gln Lys Gln Pro Lys Gly Lys His Lys Leu 180	818
aat aag ggg gtg ttt ggg gag aaa cca aac ttg cct gag tac aaa gta aat aag ggg gtg ttt ggg gag aaa cca aac ttg cct gag tac aaa gta Agn Lys Gly Val Phe Gly Glu Lys Pro Asn Leu Pro Glu Tyr Lys Val 200 200	866
gcc gcc atc cgg aag tcg ccc ttc atc ctg ttg cac tgt ggg gca ctg Ala Ala Ile Arg Lys Ser Pro Phe Ile Leu Leu His Cys Gly Ala Leu 215 220	914
aga gcc acc tgg gat ggc ttc atc ctg ctc gcc aca ctc tat gtg gct aga gcc acc tgg gat ggc ttc atc ctg ctc gcc aca ctc tat gtg gct Arg Ala Thr Trp Asp Gly Phe Ile Leu Leu Ala Thr Leu Tyr Val Ala 235	962
gtc act gtg ccc tac agc gtg tgt gtg agc aca gca cgg gag ccc agt Val Thr Val Pro Tyr Ser Val Cys Val Ser Thr Ala Arg Glu Pro Ser 245	1010
gcc gcc cgc ggc ccg ccc agc gtc tgt gac ctg gcc gtg gag gtc ctc gcc gcc cgc ggc ccg ccc agc gtc tgt gac ctg gcc gtg gag gtc ctc gc gcc cgc ggc ccg ccc agc gtc tgt gac ctg gcc gtg gag gtc ctc gc gcc gcc cgc ggc ccg ccc agc gtc tgt gac ctg gcc gtg gag gtc ctc gc gcc gcc ggc ggc ccg ccc agc gtc tgt gac ctg gcc gtg gag gtc ctc gc gcc gcc ggc ggc ccg ccc agc gtc tgt gac ctg gcc gtg gag gtc ctc gc gcc gcc ggc ggc ccg ccc agc gtc tgt gac ctg gcc gtg gag gtc ctc gc gcc gcc ggc ggc ccg ccc agc gtc tgt gac ctg gcc gtg gag gtc ctc gc gcc gcc gcc ggc ccg ccc agc gtc tgt gac ctg gcc gtg gag gtc ctc gc gcc gcc cgc ggc ccg ccc agc gtc tgt gac ctg gcc gtg gac gtc ctc gc gcc gcc gcc cgc gcc ccg ccc agc gtc tgt gac ctg gcc gtg gac ctc gc gcc gcc cgc gcc ccg ccc agc gtc tgt gac ctg gcc gtg gac ctc gc gcc gcc cgc gcc ccg ccc agc gtc tgt gac ctg gcc gcc gcc gcc gcc gcc gcc gcc gc	1058
ttc atc ctt gac att gtg ctg aat ttc cgt acc aca ttc gtg tcc aag ttc atc ctt gac att gtg ctg aat ttc cgt acc aca ttc gtg tcc aag ttc atc ctt gac att gtg ctg aat ttc cgt acc aca ttc gtg tcc aag ttc atc ctt gac att gtg ctg aat ttc cgt acc aca ttc gtg tcc aag ttc atc ctt gac att gtg ctg aat ttc cgt acc aca ttc gtg tcc aag 280 285	1106
tcg ggc cag gtg gtg ttt gcc cca aag tcc att tgc ctc cac tac gtc Ser Gly Gln Val Val Phe Ala Pro Lys Ser Ile Cys Leu His Tyr Val	1154
acc acc tgg ttc ctg ctg gat gtc atc gca gcg ctg ccc ttt gac ctg Thr Thr Trp Phe Leu Leu Asp Val Ile Ala Ala Leu Pro Phe Asp Leu 310 315	1202
cta cat gcc ttc aag gtc aac gtg tac ttc ggg gcc cat ctg ctg aag Leu His Ala Phe Lys Val Asn Val Tyr Phe Gly Ala His Leu Leu Lys 325 330	1250
acg gtg cgc ctg ctg cgc ctg ctg cgc ctg ctt ccg cgg ctg gac cgg acg gtg cgc ctg ctg cgc ctg ctg cgc ctg ctc ccg cgg ctg gac cgg acg gtg cgc ctg ctg ctg ctg ctg ctc ctg ctt ccg cgg ctg gac cgg acg gtg cgc ctg ctg ctg ctg ctg ctc ctg ctt ccg cgg ctg gac cgg acg gtg cgc ctg ctg ctg ctg ctg ctg ctc ctg ctc ccg cgg ctg gac cgg acg stg ctg ctg ctg ctg ctg ctg ctg ctg ctg c	1298
tac tcg cag tac agc gcc gtg gtg ctg aca ctg ctc atg gcc gtg ttc tac tcg cag tac agc gcc gtg gtg ctg aca ctg ctc atg gcc gtg ttc tac tcg cag tac agc gcc gtg gtg ctg aca ctg ctc atg gcc gtg ttc tac tcg cag tac agc gcc gtg gtg ctg aca ctg ctc atg gcc gtg ttc	1346
gcc ctg ctc gcg cac tgg gtc gcc tgc gtc tgg ttt tac att ggc cag gcc ctg ctc gcg cac tgg gtc gcc tgc gtc tgg ttt tac att ggc cag Ala Leu Leu Ala His Trp Val Ala Cys Val Trp Phe Tyr Ile Gly Gln 375	1394
cgg gag atc gag agc agc gaa tcc gag ctg cct gag att ggc tgg ctg Arg Glu Ile Glu Ser Ser Glu Ser Glu Leu Pro Glu Ile Gly Trp Leu 395 385	1442

cag gag ctg gcc cgc cga ctg gag act ccc tac tac ctg gtg ggc cgg Gln Glu Leu Ala Arg Arg Leu Glu Thr Pro Tyr Tyr Leu Val Gly Arg 405	90
agg cca gct gga ggg aac agc tcc ggc cag agt gac aac tgc agc agc 15 agg cca gct gga ggg aac agc tcc ggc cag agt gac aac tgc agc agc 15 agg cca gct gga ggg aac agc tcc ggc cag agt gac aac tgc agc agc 15 agg cca gct gga ggg aac agc tcc ggc cag agt gac aac tgc agc agc 15 agg cca gct gga ggg aac agc tcc ggc cag agt gac aac tgc agc agc 15 agg cca gct gga ggg aac agc tcc ggc cag agt gac aac tgc agc agc 15 agg cca gct gga ggg aac agc tcc ggc cag agt gac aac tgc agc agc 15 agg cca gct gga ggg aac agc tcc ggc cag agt gac aac tgc agc agc 15 agg cca gct gga ggg aac agc tcc ggc cag agt gac aac tgc agc agc agc agc 15	538
agc agc gag gcc aac ggg acg ggg ctg gag ctg ctg ggc ggc ccg tcg agc agc gag gcc aac ggg acg ggg ctg gag ctg ctg ggc ggc ccg tcg agc agc gag gcc aac ggg acg ggg ctg gag ctg ctg ggc ggc ccg tcg 1 agc agc gag gcc aac ggg acg ggg ctg gag ctg ctg ggc ggc ccg tcg 1 445	586
ctg cgc agc gcc tac atc acc tcc ctc tac ttc gca ctc agc agc ctc Leu Arg Ser Ala Tyr Ile Thr Ser Leu Tyr Phe Ala Leu Ser Ser Leu 455	L634
acc agc gtg ggc ttc ggc aac gtg tcc gcc aac acg gac acc gag aag acc agc gtg ggc ttc ggc aac gtg tcc gcc aac acg gac acc gag aag Thr Ser Val Gly Phe Gly Asn Val Ser Ala Asn Thr Asp Thr Glu Lys 470 475	1682
ato tto too ato tgo aco atg oto ato ggo gco otg atg cac gcg gtg ato tto too ato tgo aco atg oto ato ggo gco otg atg cac gcg gtg ato tto tco ato tgo aco atg oto ato ggo gco otg atg cac gcg ato tto tco ato tgo aco atg oto ato ggo gco otg atg cac gcg ato too too ato tgo aco atg oto ato ggo gco otg atg cac gcg ato tto tco ato tgo aco atg oto ato ggo gco otg atg cac gcg ato tto tco ato tgo aco atg otc ato ggo gco otg atg cac gcg ato tto tco ato tgo aco atg otc ato ggo gco otg atg cac gcg ato tto tco ato tgo aco atg otc ato ggo gco otg atg cac gcg ato tto tco ato tgo aco atg otc ato ggo gco otg atg cac gcg ato tto tco ato tgo aco atg otc ato ggo gco otg atg cac gcg ato tto tco ato tgo aco atg otc ato ggo gco otg atg otc ato gcg ato tto tco ato tgo aco atg otc ato ggo gco otg atg otc ato gcg ato tto tco ato tgo aco atg otc ato ggo gco otg atg otc ato gcg ato tto tco ato tgo aco atg otc ato ggo gco otg atg otc ato gcg ato tto tco ato tgo aco atg otc ato ggo gco otg atg otc ato gcg ato tto tco ato tgo aco atg otc ato gcg ato tco ato tco aco ato gcg otg atg otg at	1730
gtg ttt ggg aac gtg acg gcc atc atc cag cgc atg tac gcc cgc cgc gtg ttt ggg aac gtg acg gcc atc atc cag cgc atg tac gcc cgc cgc Val Phe Gly Asn Val Thr Ala Ile Ile Gln Arg Met Tyr Ala Arg Arg 500 500	1778
ttt ctg tac cac agc cgc acg cgc gac cag cgc gac tac atc cgc atc ttt ctg tac cac agc cgc acg cgc gac cag cgc gac tac atc cgc atc phe Leu Tyr His Ser Arg Thr Arg Asp Gln Arg Asp Tyr Ile Arg Ile 515 520	1826
cac cgt atc ccc aag ccc ctc aag cag cgc atg ctg gag tac ttc cag His Arg Ile Pro Lys Pro Leu Lys Gln Arg Met Leu Glu Tyr Phe Gln 530 535	1874
His Arg Ile Pro Lys 125 535 530 gcc acc tgg gcg gtg aac aat ggc atc gac acc acc gag ctg ctg cag gcc acc tgg gcg gtg aac aat ggc atc gac acc acc gag ctg ctg cag Ala Thr Trp Ala Val Asn Asn Gly Ile Asp Thr Thr Glu Leu Leu Gln 550 555	1922
Ala Thr Trp Ala Val ABN ABN 550 555 545 550 agc ctc cct gac gag ctg cgc gca gac atc gcc atg cac ctg cac aag ser Leu Pro Asp Glu Leu Arg Ala Asp Ile Ala Met His Leu His Lys 565 570	1970
Ser Leu Pro Asp Glu Leu Arg 5570 565 565 gag gtc ctg cag ctg cca ctg ttt gag gcg gcc agc cgc ggc tgc ctg Glu Val Leu Gln Leu Pro Leu Phe Glu Ala Ala Ser Arg Gly Cys Leu 580 580 580	2018
cgg gca ctg tct ctg gcc ctg cgg ccc gcc ttc tgc acg ccg ggc gag cgg gca ctg tct ctg gcc ctg cgg ccc gcc ttc tgc acg ccg ggc gag Arg Ala Leu Ser Leu Ala Leu Arg Pro Ala Phe Cys Thr Pro Gly 605	2066
Arg Ala Leu Ser Aca ter gtc tgc	2114
Tyr Leu lle His Gin 51 615	2162
tot ggc toc atg gag gtg ctc aag ggt ggc acc gtg ctc gcc atc cta Ser Gly Ser Met Glu Val Leu Lys Gly Gly Thr Val Leu Ala Ile Leu 635 625	

ggg aag ggc gac ctg atc ggc tgt gag ctg ccc cgg cgg gag cag gtg Gly Lys Gly Asp Leu Ile Gly Cys Glu Leu Pro Arg Arg Glu Gln Val 645	
gta aag gcc aat gcc gac gtg aag ggg ctg acg tac tgc gtc ctg cag gta aag gcc aat gcc gac gtg aag ggg ctg acg tac tgc gtc ctg cag gta aag gcc aat gcc gac gtg aag ggg ctg acg tac tgc gtc ctg cag 22 gta aag gcc aat gcc gac gtg aag ggg ctg acg tac tgc gtc ctg cag 670	258
tgt ctg cag ctg gct ggc ctg cac gac agc ctt gcg ctg tac ccc gag 2 tgt ctg cag ctg gct ggc ctg cac gac agc ctt gcg ctg tac ccc gag 2 tgt ctg cag ctg gct ggc ctg cac gac agc ctt gcg ctg tac ccc gag 2 tgt ctg cag ctg gct ggc ctg tac ccc gag 2 tgt ctg cag ctg tac ccc gag 680 685	306
ttt gcc ccg cgc ttc agt cgt ggc ctc cga ggg gag ctc agc tac aac 2 Phe Ala Pro Arg Phe Ser Arg Gly Leu Arg Gly Glu Leu Ser Tyr Asn 695 700	2354
ctg ggt gct ggg gga ggc tct gca gag gtg gac acc agc tcc ctg agc Leu Gly Ala Gly Gly Ser Ala Glu Val Asp Thr Ser Ser Leu Ser	2402
ggc gac aat acc ctt atg tcc acg ctg gag gag aag gag aca gat ggg Gly Asp Asn Thr Leu Met Ser Thr Leu Glu Glu Lys Glu Thr Asp Gly 725	2450
gag cag ggc ccc acg gtc tcc cca gcc cca gct gat gag ccc tcc agc Glu Gln Gly Pro Thr Val Ser Pro Ala Pro Ala Asp Glu Pro Ser 740 740 745	2498
ccc ctg ctg tcc cct ggc tgc acc tcc tca tcc tca gct gcc aag ctg ccc ctg ctg tcc cct ggc tgc acc tcc tca tcc tca gct gcc aag ctg ccc ctg ctg tcc cct ggc tgc acc tcc tca tcc tca gct gcc aag ctg ccc ctg ctg tcc cct ggc tgc acc tcc tca tcc tca gct gcc aag ctg ccc ctg ctg tcc cct ggc tgc acc tcc tca tcc tca gct gcc aag ctg ccc ctg ctg tcc cct ggc tgc acc tcc tca tcc tca gct gcc aag ctg ccc ctg ctg tcc cct ggc tgc acc tcc tca tcc tca fcc tca gct gcc aag ctg ccc ctg ctg tcc cct ggc tgc acc tcc tca tcc tca fcc tc	2546
cta tcc cca cgt cga aca gca ccc cgg cct cgt cta ggt ggc aga ggg cta tcc cca cgt cga aca gca ccc cgg cct cgt cta ggt ggc aga ggg Leu Ser Pro Arg Arg Thr Ala Pro Arg Pro Arg Leu Gly Gly Arg Gly 775 780	2594
agg cca ggc agg gca ggg gct ttg aag gct gag gct ggc ccc tct gct Arg Pro Gly Arg Ala Gly Ala Leu Lys Ala Glu Ala Gly Pro Ser Ala 790 795	2642
ccc cca cgg gcc cta gag ggg cta cgg ctg ccc ccc atg cca tgg aat Pro Pro Arg Ala Leu Glu Gly Leu Arg Leu Pro Pro Met Pro Trp Asn 805	2690
gtg ccc cca gat ctg agc ccc agg gta gta gat ggc att gaa gac ggc gtg ccc cca gat ctg agc ccc agg gta gta gat ggc att gaa gac ggc yal Pro Pro Asp Leu Ser Pro Arg Val Val Asp Gly 11e Glu Asp Gly 820 820	2738
tgt ggc tcg gac cag ccc aag ttc tct ttc cgc gtg ggc cag tct ggc tgt ggc tcg gac cag ccc aag ttc tct ttc cgc gtg ggc cag tct ggc tgt ggc tcg gac cag ccc aag ttc tct ttc cgc gtg ggc cag tct ggc tgt ggc tcg gac cag ccc aag ttc tct ttc cgc gtg ggc cag tct ggc tgt ggc tcg gac cag tct ggc	2786
ccg gaa tgt agc agc ccc tcc cct gga cca gag agc ggc ctg ctc Pro Glu Cys Ser Ser Pro Ser Pro Gly Pro Glu Ser Gly Leu Leu Pro Glu Cys Ser	2834
act gtt ccc cat ggg ccc agc gag gca agg aac aca gac aca ctg gac Thr Val Pro His Gly Pro Ser Glu Ala Arg Asn Thr Asp Thr Leu Asp 870 875	2882
865 870 aag ctt cgg cag gcg gtg aca gag ctg tca gag cag gtg ctg cag atg	2930

Lys Leu Arg Gln Ala Val Thr Glu Leu Ser Glu Gln Val Leu Gln Met 885 890	
880 att atc cta acq	2978
cgg gaa gga ctg cag tca ctt cgc cag gct gtg cag ctt gtc ctg gcg Arg Glu Gly Leu Gln Ser Leu Arg Gln Ala Val Gln Leu Val Leu Ala 900 905	
895	3026
Pro His Arg Git GIT 920	3074
cca gcc agc acc tcc ggg ctt ctg cag cct ctg tgt gtg gac act ggg Pro Ala Ser Thr Ser Gly Leu Leu Gln Pro Leu Cys Val Asp Thr Gly 930 935 940	
gca tcc tcc tac tgc ctg cag ccc cca gct ggc tct gtc ttg agt ggg Ala Ser Ser Tyr Cys Leu Gln Pro Pro Ala Gly Ser Val Leu Ser Gly 955	3122
act tgg ccc cac cct cgt ccg ggg cct cct ccc ctc atg gca ccc cgg Thr Trp Pro His Pro Arg Pro Gly Pro Pro Pro Leu Met Ala Pro Arg 965 970	3170
960 ccc tgg ggt ccc cca gcg tct cag agc tcc ccc tgg cct cga gcc aca ccc tgg ggt ccc cca gcg tct cag agc tcc ccc tgg cct cga gcc aca ccc tgg ggt ccc cca gcg tct cag agc tcc ccc tgg cct cga gcc aca pro Trp Gly Pro Pro Ala Ser Gln Ser Ser Pro Trp Pro Arg Ala Thr 980 980	3218
gct ttc tgg acc tcc acc tca gac tca gag ccc cct gcc tca gga gac gct ttc tgg acc tcc acc tca gac tca gag ccc cct gcc tca gga gac gct ttc tgg acc tcc acc tca gac tca gag ccc cct gcc tca gga gac gct ttc tgg acc tcc acc tca gac tca gag ccc cct gcc tca gga gac gct ttc tgg acc tcc acc tca gac tca gag ccc cct gcc tca gga gac gct ttc tgg acc tcc acc tca gac tca gag ccc cct gcc tca gga gac gct ttc tgg acc tcc acc tca gac tca gag ccc cct gcc tca gga gac gct ttc tgg acc tcc acc tca gac tca gag ccc cct gcc tca gga gac gct ttc tgg acc tcc acc tca gac tca ga	3266
ctc tgc tct gag ccc agc acc cct gcc tcc cct cct tct gag gaa ctc tgc tct gag ccc agc acc cct gcc tcc cct cct tct gag gaa Leu Cys Ser Glu Pro Ser Thr Pro Ala Ser Pro Pro Pro Ser Glu Glu 1015	3314
ggg gct agg act ggg ccc gca gag cct gtg agc cag gct gag gct acc Gly Ala Arg Thr Gly Pro Ala Glu Pro Val Ser Gln Ala Glu Ala Thr	3362
agc act gga gag ccc cca cca ggg tca ggg ggc ctg gcc ttg ccc tgg agc act gga gag ccc cca cca ggg tca ggg ggc ctg gcc ttg ccc tgg Ser Thr Gly Glu Pro Pro Pro Gly Ser Gly Gly Leu Ala Leu Pro Trp 1045	3410
gac ccc cac agc ctg gag atg gtg ctt att ggc tgc cat ggc tct ggc gac ccc cac agc ctg gag atg gtg ctt att ggc tgc cat ggc tct ggc Asp Pro His Ser Leu Glu Met Val Leu Ile Gly Cys His Gly Ser Gly Asp Pro His Ser Leu Glu Met Val Leu Ile Gly Cys His Gly Ser Gly 1060	3458
aca gtc cag tgg acc cag gaa gaa ggc aca ggg gtc t gagtaccagc Thr Val Gln Trp Thr Gln Glu Glu Gly Thr Gly Val 1075 1080	3505
cctagaacte agegttgeca ggtgtgetge catetgetgt teggeceaac eteagagt aggcagggtg gcagectece caeggaetee atgeggeeeg etggeteagg geagggae tggaagcaaa ggaggaeetg geteetgaet eteagagagg ataggetgga teeetgg aggcetetee teggeetget eetetgaeet eeeggtetee etetgeagge tggggge ggeetgagga caaggaagag etttgecate eeetgeatgt geeeetgeet etaeetg eccaaattttt atattaaaaa aaaaaataaa ataaactaaa aaaaaaaa	gcc 3625 gcc 3685 aga 3745
<210> 20	

<210> 20 <211> 1082 <212> PRT

<213> H. sapiens

Met Pro Ala Met Arg Gly Leu Leu Ala Pro Gln Asn Thr Phe Leu Asp Thr Ile Ala Thr Arg Phe Asp Gly Thr His Ser Asn Phe Val Leu Gly Asn Ala Ser Gly Gly Ala Leu Pro Val Val Tyr Cys Ser Asp Gly Phe Cys Asp Leu Thr Gly Phe Ser Arg Ala Glu Val Met Gln Arg Gly Cys Ala Cys Ser Phe Leu Tyr Gly Pro Asp Thr Ser Glu Leu Val Arg Gln Gln Ile Arg Lys Ala Leu Asp Glu His Lys Glu Phe Lys Ala Glu Leu Ile Leu Tyr Arg Lys Ser Gly Leu Pro Phe Trp Cys Leu Leu Asp Val Ile Pro Ile Lys Asn Glu Lys Gly Glu Val Ala Leu Phe Leu Val Ser 115 120 125 His Lys Asp Ile Ser Glu Thr Lys Asn Arg Gly Gly Pro Asp Arg Trp 130 135 Lys Glu Thr Gly Gly Gly Arg Arg Tyr Gly Arg Ala Arg Ser Lys 150 155 160 Gly Phe Asn Ala Asn Arg Arg Arg Ser Arg Ala Val Leu Tyr His Leu Ser Gly His Leu Gln Lys Gln Pro Lys Gly Lys His Lys Leu Asn Lys Gly Val Phe Gly Glu Lys Pro Asn Leu Pro Glu Tyr Lys Val Ala Ala 200 Ile Arg Lys Ser Pro Phe Ile Leu Leu His Cys Gly Ala Leu Arg Ala Thr Trp Asp Gly Phe Ile Leu Leu Ala Thr Leu Tyr Val Ala Val Thr Val Pro Tyr Ser Val Cys Val Ser Thr Ala Arg Glu Pro Ser Ala Ala 225 Arg Gly Pro Pro Ser Val Cys Asp Leu Ala Val Glu Val Leu Phe Ile Leu Asp Ile Val Leu Asn Phe Arg Thr Thr Phe Val Ser Lys Ser Gly Gln Val Val Phe Ala Pro Lys Ser Ile Cys Leu His Tyr Val Thr Thr Trp Phe Leu Leu Asp Val Ile Ala Ala Leu Pro Phe Asp Leu Leu His Ala Phe Lys Val Asn Val Tyr Phe Gly Ala His Leu Leu Lys Thr Val
335
325 Arg Leu Leu Arg Leu Leu Pro Arg Leu Asp Arg Tyr Ser Gln Tyr Ser Ala Val Val Leu Thr Leu Leu Met Ala Val Phe Ala Leu Leu Ala His Trp Val Ala Cys Val Trp Phe Tyr Ile Gly Gln Arg Glu 370 375 380 360 Ile Glu Ser Ser Glu Ser Glu Leu Pro Glu Ile Gly Trp Leu Gln Glu Leu Ala Arg Arg Leu Glu Thr Pro Tyr Tyr Leu Val Gly Arg Arg Pro
415 Ala Gly Gly Asn Ser Ser Gly Gln Ser Asp Asn Cys Ser Ser Ser Ser Glu Ala Asn Gly Thr Gly Leu Glu Leu Leu Gly Gly Pro Ser Leu Arg Ser Ala Tyr Ile Thr Ser Leu Tyr Phe Ala Leu Ser Ser Leu Thr Ser 455

Val Gly Phe Gly Asn Val Ser Ala Asn Thr Asp Thr Glu Lys Ile Phe Ser Ile Cys Thr Met Leu Ile Gly Ala Leu Met His Ala Val Val Phe Gly Asn Val Thr Ala Ile Ile Gln Arg Met Tyr Ala Arg Arg Phe Leu 505 Tyr His Ser Arg Thr Arg Asp Gln Arg Asp Tyr Ile Arg Ile His Arg 520 Ile Pro Lys Pro Leu Lys Gln Arg Met Leu Glu Tyr Phe Gln Ala Thr 535 -Trp Ala Val Asn Asn Gly Ile Asp Thr Thr Glu Leu Leu Gln Ser Leu Pro Asp Glu Leu Arg Ala Asp Ile Ala Met His Leu His Lys Glu Val 550 Leu Gln Leu Pro Leu Phe Glu Ala Ala Ser Arg Gly Cys Leu Arg Ala 565 585 580

Leu Ser Leu Ala Leu Arg Pro Ala Phe Cys Thr Pro Gly Glu Tyr Leu

605 580 600 Ile His Gln Gly Asp Ala Leu Gln Ala Leu Tyr Phe Val Cys Ser Gly 610 615 620 Ser Met Glu Val Leu Lys Gly Gly Thr Val Leu Ala Ile Leu Gly Lys 625 630 635 640 Gly Asp Leu Ile Gly Cys Glu Leu Pro Arg Arg Glu Gln Val Val Lys 650 Ala Asn Ala Asp Val Lys Gly Leu Thr Tyr Cys Val Leu Gln Cys Leu 660 665 670 Gln Leu Ala Gly Leu His Asp Ser Leu Ala Leu Tyr Pro Glu Phe Ala 675 680 685 Pro Arg Phe Ser Arg Gly Leu Arg Gly Glu Leu Ser Tyr Asn Leu Gly 695 Ala Gly Gly Ser Ala Glu Val Asp Thr Ser Ser Leu Ser Gly Asp 710 Asn Thr Leu Met Ser Thr Leu Glu Glu Lys Glu Thr Asp Gly Glu Gln Gly Pro Thr Val Ser Pro Ala Pro Ala Asp Glu Pro Ser Ser Pro Leu 745 Leu Ser Pro Gly Cys Thr Ser Ser Ser Ser Ala Ala Lys Leu Leu Ser 760 Pro Arg Arg Thr Ala Pro Arg Pro Arg Leu Gly Gly Arg Gly Arg Pro 775 Gly Arg Ala Gly Ala Leu Lys Ala Glu Ala Gly Pro Ser Ala Pro Pro 795 Arg Ala Leu Glu Gly Leu Arg Leu Pro Pro Met Pro Trp Asn Val Pro 810 Pro Asp Leu Ser Pro Arg Val Val Asp Gly Ile Glu Asp Gly Cys Gly 825 830 Ser Asp Gln Pro Lys Phe Ser Phe Arg Val Gly Gln Ser Gly Pro Glu 840 Cys Ser Ser Pro Ser Pro Gly Pro Glu Ser Gly Leu Leu Thr Val 855 Pro His Gly Pro Ser Glu Ala Arg Asn Thr Asp Thr Leu Asp Lys Leu 875 870 Arg Gln Ala Val Thr Glu Leu Ser Glu Gln Val Leu Gln Met Arg Glu 890 Gly Leu Gln Ser Leu Arg Gln Ala Val Gln Leu Val Leu Ala Pro His 905 Arg Glu Gly Pro Cys Pro Arg Ala Ser Gly Glu Gly Pro Cys Pro Ala Ser Thr Ser Gly Leu Leu Gln Pro Leu Cys Val Asp Thr Gly Ala Ser Ser Tyr Cys Leu Gln Pro Pro Ala Gly Ser Val Leu Ser Gly Thr Trp 935

955	
950 950 Met Ala Pro Arg Pro Trp	
945 950 955 Pro His Pro Arg Pro Gly Pro Pro Pro Leu Met Ala Pro Arg Pro Trp 975 970 965 970 Arg Ala Thr Ala Phe	
965 970 Arg Ala Thr Ala Phe Gly Pro Pro Ala Ser Gln Ser Ser Pro Trp Pro Arg Ala Thr Ala Phe 990 985 990	
985 980 Trp Thr Ser Thr Ser Asp Ser Glu Pro Pro Ala Ser Gly Asp Leu Cys 1005 1006	
995 1000 Pro Pro Ser Glu Glu Gly Ala Ser Glu Pro Ser Thr Pro Ala Ser Pro Pro Pro Pro Ser Thr	
Ser Glu Pro Ser In F10 1015 1020 1010 1010 1015 Ser Gln Ala Glu Ala Thr Ser Thr Arg Thr Gly Pro Ala Glu Pro Val Ser Gln Ala Glu Ala Thr Ser Thr 1030 1035 1035 New Arg Pro	
Arg Thr Gly Pro Ala Glu Pro Val Ser Gin Ala Glu 104	
1025 1030 Rev Cly Leu Ala Leu Pro Trp Asp Pro	
1030 1035 1025 109 Pro Pro Gly Ser Gly Gly Leu Ala Leu Pro Trp Asp Pro Gly Glu Pro Pro Pro Gly Ser Gly Gly Leu Ala Leu Pro Trp Asp Pro 1055	
His Ser Leu Glu Met Val Leu Ile Gly Cys His Gly Ser Gly Thr Val	
His Ser Leu Git Met val 1065	
1060 Gln Trp Thr Gln Glu Glu Gly Thr Gly Val 1075	
<210> 21	
<211> 1800	
<212> DNA	
<213> H. sapiens	
<220>	
<221> CDS	
.222 (346) (1057)	
<223> K+Hnov28, splice 1	
100- 21	60
<400> 21 atttgaatga ctgggttact tectagacte tteeteette tettaagtae agtatagtte atttgaatga aatetteagt etettagtte cagatgggtt etetatggta ggaatacagg tteetetgaa aatetteagt etettagtte tteeceagat etttgeeettg tagtaggttt	120
	180
	240 300
	357
acttetete ctattteet agttatatat getateatat getegetete acttetete ctattteet agttatatat getateatat getageag at gat aat gga agtteeetg aaacetggge tettgaagae geateaetgg ageag at gat aat gga agtteeetg aaacetggge tettgaagae geateaetgg ageag Asp Asp Gly	
agtttccctg aaacctgggc tools a met Asp Asia 1	
	405
gac tgg ggc tat atg atg act gac cca gtc aca tta aat gta ggt gga gac tgg ggc tat atg atg act gac cca gtc aca tta aat gta ggt gga gac tgg ggc tat atg atg act gac cca gtc aca tta aat gta ggt gga	405
gac tgg ggc tat atg atg act gac cca gtc aca tta dat gcd 35 Gly Asp Trp Gly Tyr Met Met Thr Asp Pro Val Thr Leu Asn Val Gly Gly 10 15	
Asp Trp Gly Tyl Met 100 15	
5	453
cac ttg tat aca acg tct ctc acc aca ttg acg cgt tac ccg gat tcc	
His Leu Tyr Thr Thr Ser 30	
4.3	501
atg ctt gga gct atg ttt ggg ggg gac ttc ccc aca gct cga gac cct atg ctt gga gct atg ttt ggg ggg gac ttc ccc aca gct cga gac cct	
atg ctt gga gct atg ttt ggg ggg gac ttc ccc aca gct cgc san Met Leu Gly Ala Met Phe Gly Gly Asp Phe Pro Thr Ala Arg Asp Pro	
40	549
caa ggc aat tac ttt att gat cga gat gga cct ctt ttc cga tat gtc	
caa ggc aat tac ttt att gat cga gat gga cct ctt tto og Tyr Val Gln Gly Asn Tyr Phe Ile Asp Arg Asp Gly Pro Leu Phe Arg Tyr Val	
Gln Gly Ash 1yr File 122 60	
and the gat the aag	597
ctc aac ttc tta aga act tca gaa ttg acc ttu by Leu Asp Phe Lys	
ctc aac ttc tta aga act tca gaa ttg acc tta ccg ttg sale ctc acc ttg	
70	645
gaa ttt gat ctg ctt cgg aaa gaa gca gat ttt tac cag att gag ccc gaa ttt gat ctg ctt cgg aaa gaa gca gat ttt tac cag att gag ccc	
gaa ttt gat ctg ctt cgg aaa gaa gca gat ttt tac cag abb 550 Pro Glu Phe Asp Leu Leu Arg Lys Glu Ala Asp Phe Tyr Gln Ile Glu Pro 90 95	
85	

38

ttg att cag tgt ctc aat gat cct aag cct ttg tat ccc atg gat act Leu Ile Gln Cys Leu Asn Asp Pro Lys Pro Leu Tyr Pro Met Asp Thr 110 115	93
ttt gaa gaa gtt gtg gag ctg tct agt act cgg aag ctt tct aag tac 7 ttt gaa gaa gtt gtg gag ctg tct agt act cgg aag ctt tct aag tac 7 phe Glu Glu Val Val Glu Leu Ser Ser Thr Arg Lys Leu Ser Lys Tyr 125	41
	189
gtc cat tcc tta cta gaa ggc atc tca aat tat ttt acc aag tgg aat gtc cat tcc tta cta gaa ggc atc tca aat tat ttt acc aag tgg aat Val His Ser Leu Leu Glu Gly Ile Ser Asn Tyr Phe Thr Lys Trp Asn 155 160	837
aag cac atg atg gac acc aga gac tgc cag gtt tcc ttt act ttt gga Lys His Met Met Asp Thr Arg Asp Cys Gln Val Ser Phe Thr Phe Gly 170 175	
165 CCC tgt gat tat cac cag gaa gtt tct ctt agg gtc cac ctg atg gaa CCC tgt gat tat cac cag gaa gtt tct ctt agg gtc cac ctg atg gaa CCC tgt gat tat cac cag gaa gtt tct ctt agg gtc cac ctg atg gaa CCC tgt gat tat cac cag gaa gtt tct ctt agg gtc cac ctg atg gaa 195 196 197	933
tac att aca aaa caa ggt ttc acg atc cgc aac acc cgg gtg cat cac Tyr Ile Thr Lys Gln Gly Phe Thr Ile Arg Asn Thr Arg Val His His 205	981
atg agt gag cgg gcc aat gaa aac aca gtg gag cac aac tgg act ttc atg agt gag cgg gcc aat gaa aac aca gtg gag cac aac tgg act ttc Met_Ser Glu Arg Ala Asn Glu Asn Thr Val Glu His Asn Trp Thr Phe 220 225	1029
	1077
tgt agg cta gcc cgg aag aca gac gac t gatctccgac cctgccacag	
are year len win era era era	
	1137
gttcctggaa agactctcca ggaaatggaa gatactgatt tttttttta aatcacagtg	1197
gttcctggaa agactctcca ggaaatggaa gatactgatt ttttttttta agcagaagg gaccagaagg tgagtattt tttttctttt aaatagttgt atttatttga aggcagtgag gaccagaagg tgagatattt tttttctttt aaatagttgt atttatttga ttcccctga gtatgcatgt	1257
tgagatattt tttttcccatg ttttgttccc ttcccctgg sattatggta	1317
tgagatattt ttttttttagcaga ctcctccatg ttttgttccc ttccctctga cattatggta aagttttgtg ctttggcaga ctcctccatg ttttgttccc ttccctcttggaaaat cattatggta gcctgttcag agtctccaga tacctttttt ataaaaagaa gcctgaaaaat ctataaactaa ccttaacaga gcttttctta ttacagtgct aaaataagc agagtactca tataatctac ccttaacaga gcctaaaaaat acaagaatga agagtactca aaacaataaa	1377
gcctgttcag agtttaacaga gcttttctta ttacagtgct adadgustagg agagtactca	1437
gcctgttcag agttttctta ttacagtgct aaaatgatte oggtcataatctaa ccttaacaga gcttttctta ttacagtgct aaaatgatte oggtcctaac tcaactagaa ggctaaaaat acaagaatga aagaataagc agagtactca ggtcccttac tcaactagaa ggctaaaaat gtatgccttt gagaaaaaata aaaacatcat gtagggtgac ctagtttcca aaccaataaa tgatgccttt gagaaaaaat aaaacatcat gtacaatca tttacaagtga ggaaccaaat	1497
ggtccctaat daaaaatc aaaacatcat gtagggtgat ctagtaaagta cttattaagt	1557 1617
ggtccctaac teaactagus so ggtccctaac ctagttteac tetaataagt tgatgccttt gagaaaaatc aaaacatcat gttccaatca tttaaaagta cttattaagt taagtagtat tgtaatatta aaggaaaact gttccaatca tttaaaaccaa ggaaccaaat taagtagtat tgtaatatga caactgtttc tttctatgca tatacaaacca atacaaaatg	1677
tgatgccttt gagadadattt aaggaaaact gttccaatca tttaadagta taagtagtat tgtaatatta aaggaaaact gttccaatca tttaadagta ggaaccaaat aagtagtat acagttatga caactgtttc tttctatgca tataaatcaa ggaaccaaat actgcttttt acagttatga caactgtttc tttctatga ttaaattctga atacaaaatg	1737
taagtagtat tytaatata caactgtttc tttctatgca tatadateta atacaaaatg actgcttttt acagttatga caactgtttc tttctatgca tatadateta atacaaaatg atctgtagcc atggaaatgt ctgactagaa atatttatat tgaattccca agtgtactgt atctgtagcc atggaaatgt ctctttatgc ctggtgcagt ataattccca agtgtactgt	1797
actgcttttt acagttatga etgactagaa atattatat tgaattetga etgactgt atctgtagcc atggaaatgt ctgactagaa ctggtgcagt ataattccca agtgtactgt tccctgtggt agaaaactta ctctttatgc ctggtgcagt ataattccca agtgtactgt ctaccagaaa aaaaaaacaa aactaataaa aaatgaaata tgaaaaaaaa aaaaaaaaa aaa	1800
<210> 22	
<211> 1836	
<212> DNA <213> H. sapiens	
<215> n. sep	
<220>	
-221 > CDS	
202 (202) . (1093)	
<222> (362)	

100 22	60
	120
gaggaatgtt atgatttegt agaccacatg gttgggaaag gagaaagag dataatcaca acctataget tetetetet agaccacatg gttgggaaag gagaaagag dataatcaca cataatcaca ttgtagagaa aaatccattt etgeagtggt atggttaagg gagaaagaga dataatcaca ttgtagagaa aaatccattt etgeagtggt geetgtatgt gaatttaaaa eteceaagact	180
acctatages anatocattt ctgcagtggt atggttaagg anatgttaac cccaaagact	240
acctataget tetetetee tegeagtggt atggttaagg ataatetaac eccaaagaet ttgtagagaa aaatecattt etgeagtggt geetgtatgt gaatgttaac eccaaagaet ttateettgt atgeetgget aettgtgetg geetgtatgt egettteaaa eteecacatt ttateettgt atgeetgaact agttactata aaaagtattt egetttaaac etgegetett	300 360
ttgtagagaa aaatccattt tattgtgtg gcctgtatgt gaatgttaat totccacatt ttatccttgt atgctggct acttgtgctg gcctgtatgt cgctttcaaa ctcccacatt cctttagatg tcgctgaact agttactata aaaagtattt cgctttcaaa ctgggctctt tcaagaagag caaaactcaa tacaaggcaa ttttgaagtt tccctgaaac ctgggctctt tcaagaagag caaaactcaa tacaaggcaa ttttgaagtt tccctgaaac ctgggctctt tcaagaagag caaaactcaa tacaaggcaa ttttgaagtt tccctgaaac ctgggctctt	411
	311
gaagacgcat cactggagca g atg gat aat gga gac tgg ggs Tyr Met Met Met Asp Asn Gly Asp Trp Gly Tyr Met 10	
1 5	
	459
act gac cca gtc aca tta aat gta ggt gga cac ttg tat aca acg tct act gac cca gtc aca tta aat gta ggt gga cac ttg tat aca acg tct act gac cca gtc aca tta aat gta ggt gga cac ttg tat aca acg tct act gac cca gtc aca tta aat gta ggt gga cac ttg tat aca acg tct	
act gac cca gtc aca tta aat gta ggt gga cac ttg tat act act act act act act act act act	
Thr Asp Pro val III 20	
are get atg ttt	507
acc aca ttg acg cgt tac ccg gat tcc atg ctt gg Ala Met Phe	
ctc acc aca ttg acg cgt tac ccg gat tcc atg ctt gga get deg Leu Thr Thr Leu Thr Arg Tyr Pro Asp Ser Met Leu Gly Ala Met Phe Leu Thr Thr Leu Thr Arg Tyr Pro Asp Ser Met Leu Gly Ala Met Phe	
30	555
ggg ggg gac ttc ccc aca gct cga gac cct caa ggc aat tac ttt att ggg ggg gac ttc ccc aca gct cga gac cct caa ggc aat tac ttt att	30-
ggg ggg gac ttc ccc aca gct cga gut Pro Gln Gly Asn Tyr Phe 11e	
ggg ggg gac ttc ccc aca gct cga gac cct caa ggc aat tab Phe Ile Gly Gly Asp Phe Pro Thr Ala Arg Asp Pro Gln Gly Asn Tyr Phe Ile 50 55	
45	603
gat cga gat gga cct ctt ttc cga tat gtc ctc aac ttc tta aga act	
gat cga gat gga cct ctt ttc cga tat gtc ctc aac tte tea again the first tea again the first cga tat gtc ctc aac tte tea again the gat cga gat gga gat	
Asp Arg Asp G19 1-0 65	
tca gaa ttg acc tta ccg ttg gat ttt aag gaa ttt gat ctg ctt cgg tca gaa ttg acc tta ccg ttg gat ttt aag gaa ttt gat ctg ctt cgg tca gaa ttg acc tta ccg ttg gat ttt aag gaa ttt gat ctg ctt cgg	651
tca gaa ttg acc tta ccg ttg gat ttt aag gaa ttt gat ctg belle Leu Arg Ser Glu Leu Thr Leu Pro Leu Asp Phe Lys Glu Phe Asp Leu Leu Arg 80 85	
cor Glu Leu IIII Leu - B5	
75	699
75 aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc aat aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc aat aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc aat aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc aat aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc aat aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc aat aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc aat aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc aat aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc aat aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc aat	
aaa gaa gca gat ttt tac tag abo glu Pro Leu Ile Gln Cys Leu Aba	
aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tge bei Asn Lys Glu Ala Asp Phe Tyr Gln Ile Glu Pro Leu Ile Gln Cys Leu Asn 100 105	
are gar	747
and sag cot ttg tat coc atg gat act ttt gad gat val Val Glu	
gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gas gat	
ASP PIO 210 115	795
ctg tot agt act cgg aag oft tot aag tac too aac cca gtg got gto	,,,,
ctg tct agt act cgg aag ctt tct duy Tyr Ser Asn Pro Val Ala Val	
ctg tct agt act cgg aag ctt tct aag tac tcc aac cca geg solval val Leu Ser Ser Thr Arg Lys Leu Ser Lys Tyr Ser Asn Pro Val Ala Val 130	
125	843
atc ata acg caa cta acc atc acc act aag gtc cat tcc tta cta gaa atc ata acg caa cta acc atc acc act aag gtc cat tcc tta cta gaa atc ata acg caa cta acc atc acc act aag gtc cat tcc tta cta gaa	
atc ata acg caa cta acc atc acc act aag gtc cat tee tee our Glu Ile Ile Thr Gln Leu Thr Ile Thr Thr Lys Val His Ser Leu Leu Glu 145	
Ile lie illi Gin 254 145	001
ggc atc tca aat tat ttt acc aag tgg aat aag cac atg atg gac acc ggc atc tca aat tat ttt ttt acc aag tgg aat aag cac atg atg gac acc	891
ggc atc tca aat tat ttt acc aag tgg aat aag cac atg acg asp Thr Gly Ile Ser Asn Tyr Phe Thr Lys Trp Asn Lys His Met Met Asp Thr 160 165	
Cly Ile Ser Ash 191 225 165	
155	939
hat fat cac cag	
aga gac tgc cag gtt tee ttt and the gly pro Cys Asp Tyr His old	
Arg Asp Cys Gin value 180	
173 aaa caa qgt	987
gaa gtt tct ctt agg gtc cac ctg atg gaa tac att aca aaa caa ggt	
gaa gtt tct ctt agg gtc cac ctg atg gaa tac att aca tac Glu Glu Glu Val Ser Leu Arg Val His Leu Met Glu Tyr Ile Thr Lys Gln Gly	
190	1035
ttc acg atc cgc aac acc cgg gtg cat cac atg agt gag cgg gcc aat ttc acg atc cgc aac acc cgg gtg cat cac atg agt gag cgg gcc aat	
ttc acg atc cgc aac acc cgg gtg cat cac atg agt gag cgs street tc acg atc cgc aac acc cgg gtg cat cac atg agt gag cgs street acg atc acg atg agt gag cgs street acg atc acg agt gag cgs street acg atc acg agt gag cgs street	
Phe Thr Ile Arg Asn Thr Arg var	
3V .	

215		
205 210 cm tot agg cta gcc cq	g aag	.083
gaa aac aca gtg gag cac aac tgg act ttc tgt agg cta gcc cg Glu Asn Thr Val Glu His Asn Trp Thr Phe Cys Arg Leu Ala Ar	g Lys	
220		1133
aca gac gac t gateteegac cetgecacag gtteetggaa agacteteea Thr Asp Asp 235		
gatactgatt tttttttta aatcacagtg tgagatattt tt	ttggcaga	1193 1253
aaatagttgt atttattee tteeceetga gtatgeatgt geetgtteag ag	ttaacaga	1313 1373
		1433 1493
		1553 1613
aaaacatcat gtagggtgaa oosaa	tggaaatgt	1673
		1733 1793
ctgactagaa atatttataa ataattagaa agtgtactgt ctaccagaaa a	auauau	1836
ctctttatgc ctggtgcagt atdattccod eg.gc aactaataaa aaatgaaata tgaaaaaaaaa aaaaaaaaa		
<210> 23 <211> 1751		
<212> DNA		
<213> H. sapiens		
<220>		
<221> CDS <222> (297)(1008)		
<223> K+Hnov28 splice 3		
<400> 23 ccatgtttct taccatgtct tgccagagct ttagaaattt gctctgcagt ccatgtttct taccatgtct tgccagagg gaactgacta aggcagttca	ttgctttaca	60 120
		180 240
		299
gtatetgage attteteagt gtettaagge tggeteteda tgagtgetgg gtatetgage attteteagt gtettaagge etettgaaga egeateaetg eteatetata tegttteeet gaaacetggg etettgaaga egeateaetg	Met 1	
-t 20	a rra aat	347
gat aat gga gac tgg ggc tat atg atg act gac cca gtc ac Asp Asn Gly Asp Trp Gly Tyr Met Met Thr Asp Pro Val Th	r Leu Asn	
y and the ac	o cqt tac	395
gta ggt gga cac ttg tat aca acg tct ctc acc aca ttg ac Val Gly Gly His Leu Tyr Thr Thr Ser Leu Thr Thr Leu Th 20 25 30		
and dag tto C	cc aca gct	443
pro Asp Ser Met Lea 9-1 45		491
cga gac cct caa ggc aat tac ttt att gat cga gat gga c	ct ctt ttc ro Leu Phe	471
Arg Asp Pro Gin Gly 7555 60	65	
50	ta ccg ttg	539
cga tat gtc ctc aac ttc tta aga act tca gaa ttg acc t Arg Tyr Val Leu Asn Phe Leu Arg Thr Ser Glu Leu Thr I 70	Leu Pro Leu 80	

WO 99/43696

gat ttt aag gaa ttt gat ctg ctt cgg aaa gaa gca gat ttt tac cag Asp Phe Lys Glu Phe Asp Leu Leu Arg Lys Glu Ala Asp Phe Tyr Gln 85 90 95	587
	635
atg gat act ttt gaa gaa gtt gtg gag ctg tct agt act cgg aag ctt Met Asp Thr Phe Glu Glu Val Val Glu Leu Ser Ser Thr Arg Lys Leu 120 125	683
tct aag tac tcc aac cca gtg gct gtc atc ata acg caa cta acc atc tct aag tac tcc aac cca gtg gct gtc atc ata acg caa cta acc atc tct aag tac tcc aac cca gtg gct gtc atc ata acg caa cta acc atc tct aag tac tcc aac cca gtg gct gtc atc ata acg caa cta acc atc tct aag tac tcc aac cca gtg gct gtc atc ata acg caa cta acc atc 140 140 145	731
acc act aag gtc cat tcc tta cta gaa ggc atc tca aat tat ttt acc acc act aag gtc cat tcc tta cta gaa ggc atc tca aat tat ttt acc Thr Thr Lys Val His Ser Leu Leu Glu Gly Ile Ser Asn Tyr Phe Thr 150 150 160	779
aag tgg aat aag cac atg atg gac acc aga gac tgc cag gtt tcc ttt Lys Trp Asn Lys His Met Met Asp Thr Arg Asp Cys Gln Val Ser Phe 165 170 175	827
act ttt gga ccc tgt gat tat cac cag gaa gtt tct ctt agg gtc cac Thr Phe Gly Pro Cys Asp Tyr His Gln Glu Val Ser Leu Arg Val His	875
ctg atg gaa tac att aca aaa caa ggt ttc acg atc cgc aac acc cgg Leu Met Glu Tyr Ile Thr Lys Gln Gly Phe Thr Ile Arg Asn Thr Arg 200 205	923
gtg cat cac atg agt gag cgg gcc aat gaa aac aca gtg gag cac aac gtg cat cac atg agt gag cgg gcc aat gaa aac aca gtg gag cac aac Val His His Met Ser Glu Arg Ala Asn Glu Asn Thr Val Glu His Asn 215 220 225	971
tgg act ttc tgt agg cta gcc cgg aag aca gac gac t gatctccgac tgg act ttc tgt agg cta gcc cgg aag aca gac gac t gatctccgac Trp Thr Phe Cys Arg Leu Ala Arg Lys Thr Asp Asp 230 235	1018
cctgccacag gttcctggaa agactctcca ggaaatggaa gatactgatt tttttttta aaatagatggaa ggaaatggaa ggaaatggag aggcagtgag gaccagaagg gagttttgtg ctttggcaga gtctccaga gcctgttcag agtctccaga gcttttctta taaaaaaaa gcctgataaaat ggccctaac cctaacagaa gctttctta taaactaaa ggccctaac tcaactagaa ggcagtagcat taaaatcaa aggaaaaatc aggaaaaaatc tgaagcatt tgaaaatata accaataaa cttataagt accgctttt aagaaaaatc aaaggaaaaact accaacaaaa accgcttttt aagaaaaacta ggaaaaacta ggaaaaacta ggaaaaacta ggaaaaacta ccaactagaa accggaaaaat ccaactagaa accggaaaaacta accggaaaaacta accggaaaaacta accggaaaaacta accggaaaaacta ccaactagaa accggaaaaaaaaaa	10,0
<pre><210> 24 <211> 1542 <212> DNA <213> H. sapiens</pre>	

<220>

<221> CDS

<221> CDS <222> (88)(799)	
<222> (88)(1997) <223> K+Hnov28, splice 4	
<pre><400> 24 cgggcatctc ccggcccggc cgcagcagcc gccgccgccg cgcatttccc tgaaacctgg cgggcatctc ccggcccggc cgcagcagcag atg gat aat gga gac tgg ggc tat atg gctcttgaag acgcatcact ggaagcag atg gat aat gga gac tgg ggc tat atg Met Asp Asn Gly Asp Trp Gly Tyr Met 1 5</pre>	60 114
atg act gac cca gtc aca tta aat gta ggt gga cac ttg tat aca acg atg act gac cca gtc aca tta aat gta ggt gga cac ttg tat aca acg atg act gac cca gtc aca tta aat gta ggt gga cac ttg tat aca acg atg act gac cca gtc aca tta aat gta ggt gga cac ttg tat aca acg atg act gac cca gtc aca tta aat gta ggt gga cac ttg tat aca acg 25	162
tct ctc acc aca ttg acg cgt tac ccg gat tcc atg ctt gga gct atg tct ctc acc aca ttg acg cgt tac ccg gat tcc atg ctt gga gct atg tct ctc acc aca ttg acg cgt tac ccg gat tcc atg ctt gga gct atg tct ctc acc aca ttg acg cgt tac ccg gat tcc atg ctt gga gct atg tct ctc acc aca ttg acg cgt tac ccg gat tcc atg ctt gga gct atg tct ctc acc aca ttg acg cgt tac ccg gat tcc atg ctt gga gct atg tct ctc acc aca ttg acg cgt tac ccg gat tcc atg ctt gga gct atg tct ctc acc aca ttg acg cgt tac ccg gat tcc atg ctt gga gct atg	210
ttt ggg ggg gac ttc ccc aca gct cga gac cct caa ggc aat tac ttt ttt ggg ggg gac ttc ccc aca gct cga gac cct caa ggc aat tac ttt phe Gly Gly Asp Phe Pro Thr Ala Arg Asp Pro Gln Gly Asn Tyr Phe 50 55	258
att gat cga gat gga cct ctt ttc cga tat gtc ctc aac ttc tta aga att gat cga gat gga cct ctt ttc cga tat gtc ctc aac ttc tta aga Ile Asp Arg Asp Gly Pro Leu Phe Arg Tyr Val Leu Asn Phe Leu Arg 65	306
act tca gaa ttg acc tta ccg ttg gat ttt aag gaa ttt gat ctg ctt Thr Ser Glu Leu Thr Leu Pro Leu Asp Phe Lys Glu Phe Asp Leu Leu 80 85	354
cgg aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc cgg aaa gaa gca gat ttt tac cag att gag ccc ttg att cag tgt ctc Leu Arg Lys Glu Ala Asp Phe Tyr Gln Ile Glu Pro Leu Ile Gln Cys Leu 105	402
aat gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gtg aat gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gtg aat gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gtg aat gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gtg aat gat gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gtg aat gat gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gtg aat gat gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gtg aat gat gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gtg aat gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gtg aat gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gtg aat gat gat cct aag cct ttg tat ccc atg gat act ttt gaa gaa gtt gtg aat gat gat gat gat gat gat gat gat gat	450
gag ctg tct agt act cgg aag ctt tct aag tac tcc aac cca gtg gct Glu Leu Ser Ser Thr Arg Lys Leu Ser Lys Tyr Ser Asn Pro Val Ala 130 135	498
gtc atc ata acg caa cta acc atc acc act aag gtc cat tcc tta cta Val Ile Ile Thr Gln Leu Thr Ile Thr Thr Lys Val His Ser Leu Leu 145	546
gaa ggc atc tca aat tat ttt acc aag tgg aat aag cac atg atg gac glu Gly Ile Ser Asn Tyr Phe Thr Lys Trp Asn Lys His Met Met Asp 160 165	594
acc aga gac tgc cag gtt tcc ttt act ttt gga ccc tgt gat tat cac acc aga gac tgc cag gtt tcc ttt act ttt gga ccc tgt gat tat cac Thr Arg Asp Cys Gln Val Ser Phe Thr Phe Gly Pro Cys Asp Tyr His 185	642
cag gaa gtt tot ott agg gto cac otg atg gaa tac att aca aaa caa cal gaa gtt tot ott agg gto cac otg atg gaa tac att aca aaa caa cag gaa gtt tot ott agg gto cac otg atg gaa tac att aca aaa caa cag gaa gtt tot ott agg gto cac otg atg gaa tac att aca aaa caa cag gaa gtt tot ott agg gto cac otg atg gaa tac att aca aaa caa cag gaa gtt tot ott agg gto cac otg atg gaa tac att aca aaa caa cag gaa gtt tot ott agg gto cac otg atg gaa tac att aca aaa caa cag gaa gtt tot ott agg gto cac otg atg gaa tac att aca aaa caa cag gaa gtt tot ott agg gto cac otg atg gaa tac att aca aaa caa cag gaa gto otg otg otg otg otg otg otg otg otg	690
ggt ttc acg atc cgc aac acc cgg gtg cat cac atg agt gag cgg gcc Gly Phe Thr Ile Arg Asn Thr Arg Val His His Met Ser Glu Arg Ala 215 205 43	738

aat gaa aac aca gtg gag cac aac tgg act ttc tgt agg cta gcc cgg Asn Glu Asn Thr Val Glu His Asn Trp Thr Phe Cys Arg Leu Ala Arg 225 230	786
aag aca gac gac t gateteegae eetgecacag gtteetggaa agaeteteea	839
Lys Thr Asp Asp	899
therefore a attacacagts transaction to the contract to the con	959
235 ggaaatggaa gatactgatt ttttttttta aatcacagtg tgagatattt tttttctttt ggaaatggaa gatactgatt ttttttttta aatcacagtg tgagatatttgtg ctttggcaga ggaaatggaa gatactgattag aggcagtgag gaccagaagg aagttttgg agtctccaga	1019
aaatagttgt atttattagg treccettga gtatgeatgt geetgetatag gettaacaga	1079
ctcctccatg tittiggaaaat cattatggta tataactagaa tgaactagaa	1139
ctcctccatg ttttgtttct tataaaaagaa gtctgaaaaat cattatggta tataatctat ttacattagta tacctttttt ataaaaagaa gtctgaaaaat ctgataaaat ggtccctaac tcaactagaa gcttttctta ttacagtgct aaaatgattt ctgataaaaat gagtactca tgatgccttt gagaaaaatc ggctaaaaaat acaagaatga aagaataagc agagtactca taagtagtat tgtaatatta ggctaaaaaat acaggtagc ctagtttcca aaccaataaa taagtagtat tgtaatatga	1199
gcttttctta ttatagss	1259
gettetetta ttacagogeta aagaataage agagtaetea tgatgetete gegetaatatta ggetaaaaat acaagaatga aagaataage aaccaataaa taagtagtat tgtaatatta aaaccateat gtagggtgae etagteteea aaccaataaa taagtagtat tacagtetet acagtetatga aaaacateat gtaggagtga tttaaaaagta ettattaagt aetgetettt acagttatga	1319 1379
ggctaaaaat acaagaatga ctagtttcca aaccaataaa taagtagtat agattatga aaacatcat gtagggtgac ctagtttcca atctattaagt actgcttttt acagttatga aaggaaaact gttccaatca tttaaaagta cttattaagt atctgtagcc atggaaatgt atgaaaactaa ggaaccaaat atctgtagcc atggaaactta	1439
aaacatcat gtagggegaa tttaaaaagta cttattaagt actgettte atggaaatgt aaggaaaact gttccaatca tttaaaatcaa ggaaccaaat atctgtagcc atggaaactta caactgttte tttctatgca tataaatcaa atacaaaatg tcccttgtggt agaaaactta caactgttte tttctatat tgaattctga atacaaaatg tcccttgtaga aaaaaaacaa	1499
atgactagaa didicious sangtatactat claccaguan	1542
caactgtttc tttctatgda tgaattctga atacaaaatg tccctgtggt dgaacaacaa ctgactagaa atatttatat tgaattccca agtgtactgt ctaccagaaa aaaaaaaacaa ctctttatgc ctggtgcagt ataattccca agtgtactgt ctaccagaaa aaaaacaaa ctctttatgc ctgaaata tgaaaaaaaaa aaaaaaaaaa	
ctctttatgc ctggtgcagt ataattccca agtguers aactaataaa aaatgaaata tgaaaaaaaa aaaaaaaaaa	
<210> 25 <211> 237	
<211> 25 PRT	
<213> H. sapiens	
<400> 25 Cly Tyr Met Met Thr Asp Pro Val IIII 150	
<pre><400> 25 Met Asp Asn Gly Asp Trp Gly Tyr Met Met Thr Asp Pro Val Thr Leu 15 10</pre>	
1 Wie Leu Tyr Thr Thr Ser beu im 30	
Asn Val Gly Gly Ars 25 20 20 Tyr Pro Asp Ser Met Leu Gly Ala Met Phe Gly Gly Asp Phe Pro Thr 45 40 Asn Day Asp Gly Pro Leu	
Tyr Pro Asp Ser Met Leu Gly Ald Met 115 45	
Tyr Pro Asp Ser Met 201 40 35 40 Ala Arg Asp Pro Gln Gly Asn Tyr Phe Ile Asp Arg Asp Gly Pro Leu 55 60 Ala Arg Asp Pro Gln Gly Asn Tyr Phe Ile Asp Arg Asp Gly Pro Leu 60 55 60 60 60 60 60 60 60 60 60 60 60 60 60	
Ala Arg Asp Pro Gin Gif 55 60	
Ala Arg Asp Pro Gill 55 50 50 Phe Arg Tyr Val Leu Asn Phe Leu Arg Thr Ser Glu Leu Thr Leu Pro 70 70 70 70 71 72 73 74 75 75 76 77 78 78 79 70 70 70 70 70 70 70 70 70	
phe Arg Tyl Var Tyr 70 70 Tys Glu Ala Asp Phe Tyr	
Phe Arg Tyr Val Let 75 70 65 Leu Asp Phe Lys Glu Phe Asp Leu Leu Arg Lys Glu Ala Asp Phe Tyr 95 100 100 100 100 100 100 100	
85 Cyr Cyr Leu Asn Asp Pro Lys Pro Leu Tyl	
Gln Ile Glu Pro Leu Ile Gln Cys Leu Asn Asp Pro Lys Pro Leu Tyr	
The Dhe Clu Glu Val Val Glu Leu Ser 325	
120 115 120 125 Thr Gln Leu Thr Leu Ser Lys Tyr Ser Asn Pro Val Ala Val Ile Ile Thr Gln Leu Thr 140 135 140 Ser Asn Tyr Phe	
Cor Lys Tyr Ser Asn Pro Val Ala Val He 110 140	
Leu Ser Lys Tyl Set 135 135 136 137 138 138 139 140 140 140 140 140 140 140 14	
The Thr Lys Val His Ser Leu neu 014 155	
150 145 Thr Arg Asp Cys Gln Val Ser	
The Thr Lys Val 155 150 150 150 155 145 Thr Lys Trp Asn Lys His Met Met Asp Thr Arg Asp Cys Gln Val Ser 175 165 Phe Thr Phe Gly Pro Cys Asp Tyr His Gln Glu Val Ser Leu Arg Val 190 185	
phe Thr Phe Gly Pro Cys Asp Tyr HIS GIM 5190	
180 The Lys Gln Gly Phe Thr Ile Arg Ash Inc	
Phe Thr Phe Gly Flo 67 185 185 180 180 His Leu Met Glu Tyr Ile Thr Lys Gln Gly Phe Thr Ile Arg Asn Thr 205 200 200 Thr Val Glu His	;
His Leu Met Glu Tyr 200 200 195 200 195 Arg Val His His Met Ser Glu Arg Ala Asn Glu Asn Thr Val Glu His 220 220 215	
Arg Val His His Met 3215 220	
215 210 Asn Trp Thr Phe Cys Arg Leu Ala Arg Lys Thr Asp Asp 235 230 235	
225	
<210> 26	

<211> 3204

<212> DNA	
<213> H. sapiens	
<220>	
-221 > CDS	
<222> (182) · · · (1349)	
<223> K+Hnov42	
AAAA 26 tatatagaa agggaaggaa	60
<400> 26 cggccgaacc ttgggtgtgg gacagagtgc gtgcgtgtgg tgtgtcccca agggcaggaa cggccgaacc ttgggtgtgg gacagagtgg ggaggaggg gaagggcggg aggagaaaaa	120
	180 229
ggtgggagga ggacteagard ttc ctg aac ggc agc ccc aag ddo 33 g atg agg cgg gtg acc ctg ttc ctg aac ggc agc ccc aag ddo 33 g atg agg cgg gtg acc ctg ttc ctg aac ggc agc ccc aag ddo 33 Met Arg Arg Val Thr Leu Phe Leu Asn Gly Ser Pro Lys Asn Gly Lys 10	
	277
tet ata acc age	211
gtg gtt gct gta tat gga act tta tct gat ttg ctt tct ges ges gtg gtt gct tct ges ges gtg gtg gtt gct tct ges ges gtg gtg gtt gct gct tct ges ges gtg gtg gtt gct tct ges ges gtg gtg gtg gtg gtg gtg gtg gtg gtg gt	
agt aaa ctc ggc ata aaa gcc acc agt gtg tat aat ggg aaa ggt gga	325
agt aaa ctc ggc ata aaa gcc acc agt gtg tat aat ggg taa 35 56 Ser Lys Leu Gly Ile Lys Ala Thr Ser Val Tyr Asn Gly Lys Gly Gly	
Ser Lys Leu Gly 11e 272 40	
att tra ttt qtt	373
ctg att gat gat att gct ttg atc agg gat gat gat gtt teg phe Val Leu Ile Asp Asp Ile Ala Leu Ile Arg Asp Asp Asp Val Leu Phe Val	
Leu Ile Asp Asp IIe	
50	421
tgt gaa gga gag cca ttt att gat cct cag aca gat tct aag cct cct tgt gaa gga gag cca ttt att gat cct cag aca gat tct aag cct cct tgt gaa gga gag cca ttt att gat cct cag aca gat tct aag cct cct Tgt gaa gga gag cca ttt att gat cct cag aca gat tct aag cct cct tgt gaa gga gag cca ttt att gat cct cag aca gat tct aag cct cct Tgt gaa gga gag cca ttt att gat cct cag aca gat tct aag cct cct Tgt gaa gga gag cca ttt att gat cct cag aca gat tct aag cct cct Tgt gaa gga gag cca ttt att gat cct cag aca gat tct aag cct cct Tgt gaa gga gag cca ttt att gat cct cag aca gat tct aag cct cct Tgt gaa gga gag cca ttt att gat cct cag aca gat tct aag cct cct Tgt gaa gga gag cca ttt att gat cct cag aca gat tct aag cct cct	
Cys Glu Gly Glu Pro Phe 116 ASP 75	
65	469
gag gga ttg tta gga ttc cac aca gac tgg ctg aca tta aat gtt gga gag gga ttg tta gga ttc cac aca gac tgg ctg aca tta aat gtt gga	
gag gga ttg tta gga ttc cac aca gac tgg ctg aca tta dab yal Gly Glu Gly Leu Leu Gly Phe His Thr Asp Trp Leu Thr Leu Asn Val Gly 95	
os ana gaa cct	517
ggg cgg tac ttt aca act aca cgg agc act tta gtg aat aaa gaa cct ggg cgg tac ttt aca act aca cgg agc act tta gtg aat aaa gaa cct ggg cgg tac ttt aca act aca cgg agc act tta gtg aat aaa gaa cct	
Gly Arg Tyl File 112 105	
100	565
gac agt atg ctg gcc cac atg ttt aag gac aaa ggt gtc tgg gga aat	
gac agt atg ctg gcc cac atg ttt aag gac aaa ggt gcc sas 333 Asp Ser Met Leu Ala His Met Phe Lys Asp Lys Gly Val Trp Gly Asn 120	
	613
aag caa gat cat aga gga gct ttc tta att gac cga agt cct gag tac aag caa gat cat aga gga gct ttc tta att gac cga agt cct gag tac	
two Gin ASD nis Arg 4-1	
	661
ttc gaa ccc att ttg aac tac ttg cgt cat gga cag ctc att gta aat ttc gaa ccc att ttg aac tac ttg cgt cat gga cag ctc att gta aat	
ttc gaa ccc att ttg aac tac ttg cgt cat gga cag ctc atb 34 Asn Phe Glu Pro Ile Leu Asn Tyr Leu Arg His Gly Gln Leu Ile Val Asn 150 155	
	709
gat ggc att aat tta ttg ggt gtg tta gaa gaa gca aga ttt ttt ggt gat ggc att aat tta ttg ggt gtg tta gaa gaa gca aga ttt ttt ggt gat ggc att aat tta ttg ggt gtg tta gaa gaa gca aga ttt ttt ggt	
gat ggc att aat tta ttg ggt gtg tta gaa gaa gca agu bhe Phe Gly Asp Gly Ile Asn Leu Leu Gly Val Leu Glu Glu Ala Arg Phe Phe Gly	
	757
att gac tca ttg att gaa cac cta gaa gtg gca ata aag aat tct caa att gac tca ttg att gaa cac cta gaa gtg gca ata aag aat tct caa	,,,,
att gac tca ttg att gaa cac cta gaa gtg gca ata aag dab ser Gln Ile Asp Ser Leu Ile Glu His Leu Glu Val Ala Ile Lys Asn Ser Gln 185	
	805
cca ccg gag gat cat tca cca ata tcc cga aag gaa ttt gtc cga ttt	603
cca ccg gag gat cat tca cca ded 555 5	

Pro Pro Glu Asp His Ser Pro Ile Ser Arg Lys Glu Phe Val Arg Phe 200 205	
ttg cta gca act cca acc aag tca gaa ctg cga tgc cag ggt ttg aac ttg cta gca act cca acc aag tca gaa ctg cga tgc cag ggt ttg aac Leu Leu Ala Thr Pro Thr Lys Ser Glu Leu Arg Cys Gln Gly Leu Asn 215 220	853
ttc agt ggt gct gat ctt tct cgt ttg gac ctt cga tac att aac ttc ttc agt ggt gct gat ctt tct cgt ttg gac ctt cga tac att aac ttc ttc agt ggt gct gat ctt tct cgt ttg gac ctt cga tac att aac ttc ttc agt ggt gct gat ctt tct cgt ttg gac ctt cga tac att aac ttc 230	901
aaa atg gcc aat tta agc cgc tgt aat ctt gca cat gca aat ctt tgc aaa atg gcc aat tta agc cgc tgt aat ctt gca cat gca aat ctt tgc Lys Met Ala Asn Leu Ser Arg Cys Asn Leu Ala His Ala Asn Leu Cys Lys Met Ala Asn Leu Ser Arg Cys Asn Leu Ala His Ala Asn Leu Cys 250 255	949
tgt gca aat ctt gaa cga gct gat ctc tct gga tca gtg ctt gac tgt tgt gca aat ctt gaa cga gct gat ctc tct gga tca gtg ctt gac tgt Cys Ala Asn Leu Glu Arg Ala Asp Leu Ser Gly Ser Val Leu Asp Cys Cys Ala Asn Leu Glu Arg Ala Asp 265	997
gcg aat ctc cag gga gtc aag atg ctc tgt tct aat gca gaa gga gca Ala Asn Leu Gln Gly Val Lys Met Leu Cys Ser Asn Ala Glu Gly Ala	1045
tcc ctg aaa ctg tgt aat ttt gag gat cct tct ggt ctt aaa gcc aat Ser Leu Lys Leu Cys Asn Phe Glu Asp Pro Ser Gly Leu Lys Ala Asn 295 300	1093
tta gaa ggt gct aat ctg aaa ggt gtg gat atg gaa gga agt cag atg tta gaa ggt gct aat ctg aaa ggt gtg gat atg gaa gga agt cag atg Leu Glu Gly Ala Asn Leu Lys Gly Val Asp Met Glu Gly Ser Gln Met Leu Glu Gly Ala Asn Leu Lys Gly Val Asp 315	1141
aca gga att aac ctg aga gtg gct acc tta aaa aat gca aag ttg aag aca gga att aac ctg aga gtg gct acc tta aaa aat gca aag ttg aag Thr Gly Ile Asn Leu Arg Val Ala Thr Leu Lys Asn Ala Lys Leu Lys 335 336	1189
aac tgt aac ctc aga gga gca act ctg gca gga act gat tta gag aat Asn Cys Asn Leu Arg Gly Ala Thr Leu Ala Gly Thr Asp Leu Glu Asn 345	1237
tgt gat ctg tct ggg tgt gat ctt caa gaa gcc aac ctg aga ggg tcc tgt gat ctg tct ggg tgt gat ctt caa gaa gcc aac ctg aga ggg tcc Cys Asp Leu Ser Gly Cys Asp Leu Gln Glu Ala Asn Leu Arg Gly Ser 360 365	1285
aac gtg aag gga gct ata ttt gaa gag atg ctg aca cca cta cac atg aac gtg aag gga gct ata ttt gaa gag atg ctg aca cca cta cac atg acc yal Lys Gly Ala Ile Phe Glu Glu Met Leu Thr Pro Leu His Met	1333
375 370 375 tca caa agt gtc aga t gagaatttta ggggctggag gaagatgtaa aagatgaaa: Ser Gln Ser Val Arg	a 1389
385	t 1449
tgttttcctt atcacttttc tttctccacc cactcagttg tctagaagaa ataacactg tgttttcctt atcacttttc tttctccacc cactcagttg ttttgagtgg tgcataagg aaggaaattt taaaaaaaaaa catttagagg attatgcttg ttttaacaga aaagcactca tttaataga	g 1509
aaggaaattt taaaaaaddaa cuttuusissi ttttaacaga aaagcactca tttdatay	g 1629
ALSTCLACCO COCCO	
SUBTILICION WOWS	
accgtatgaa tatggtgaga tcagactccc taagactctt tttaggttaa tatccagttc ttactgat gtttactttt taggacagaa cagtagctaa attaaagtaa tatccagttc ttactgat	

agacagagtg gaaagaaaga catcattgta catcactgtc attccaaagg tacagtgtaa 2049 ctctggatgg aggaataact tacctatcac tacaacactt acaaatgaga atttctcaga 2109 atttcattct aggcaagttc cactcaacac cagatcaagc aattctatct atttacacta 2169 2229 ttagcctagt tttctcatac agtcatcaca agcataggaa gatacttcaa aaccaaaaaa accaaggtgc atcattaata ttcatttaat tcaaatacca aatagtttac atagggccag 2289 cttagaaata gatactaaat ccagagctac tgcaatcaaa gcttatatga gtgaatatgg 2349 tagagttgcc tgctaaaagg caatgtaata taattgcagc tagaacccta cagtggggaa tgaggaattt taaacacaca tttgattaca gccaccaaaa aaatagacgt aaaaataaag 2469 gcatttggct ggtccaagat gtaattttca atcagtcagc acctgtgatt cttttactta 2529 tttttttgtg gtttttttt tttaaacaaa ttttagccca attttcttga gtcattctct 2589 ctctgcagca gcagaggaag ggcctgtacc tccctaccaa tgacttggtg tccttatttt ctaccccaag agcagggata ttagctgtgt ccaaatgggt tctgaattct acagactcat 2709 caacatgagg caaggaatca ttgaaaacca cctgtgtctc ctttgggaga atgacatatc tttagtattt acgtagctta ttcttctata tctacatatg caaagctttc cttaacagta 2829 aagggtacat atgcatagtg ggaggagatc agacctttac aagtgaagga aagcaacttc agaaatgaat tattttcttt gctttattat ttttaccaag acagagaagt attgtattga 2949 gagataatct attttcataa tcaatatgtg cctaaattat atttaaatca tttcactctg 3009 tactatattt tcaggaatta cagaatgtgg tattcattca cttaaaggta cctctgtaga 3069 aataacctaa aactgcagaa ggatctgaaa gatctaaaca tggtgtgctt agaaactgca 3129 3189 gattttagat ctaatgtata ctgcattaat aaatgatata aagtgtttgt tgaaaaaaaa 3204 aaaaaaaaaa aaaaa <210> 27

<211> 389 <212> PRT <213> H. sapiens

Met Arg Arg Val Thr Leu Phe Leu Asn Gly Ser Pro Lys Asn Gly Lys_ Val Val Ala Val Tyr Gly Thr Leu Ser Asp Leu Leu Ser Val Ala Ser Ser Lys Leu Gly Ile Lys Ala Thr Ser Val Tyr Asn Gly Lys Gly Gly Leu Ile Asp Asp Ile Ala Leu Ile Arg Asp Asp Val Leu Phe Val Cys Glu Gly Glu Pro Phe Ile Asp Pro Gln Thr Asp Ser Lys Pro Pro Glu Gly Leu Leu Gly Phe His Thr Asp Trp Leu Thr Leu Asn Val Gly Gly Arg Tyr Phe Thr Thr Thr Arg Ser Thr Leu Val Asn Lys Glu Pro Asp Ser Met Leu Ala His Met Phe Lys Asp Lys Gly Val Trp Gly Asn Lys Gln Asp His Arg Gly Ala Phe Leu Ile Asp Arg Ser Pro Glu Tyr Phe Glu Pro Ile Leu Asn Tyr Leu Arg His Gly Gln Leu Ile Val Asn Asp Gly Ile Asn Leu Leu Gly Val Leu Glu Glu Ala Arg Phe Phe Gly Ile Asp Ser Leu Ile Glu His Leu Glu Val Ala Ile Lys Asn Ser Gln Pro Pro Glu Asp His Ser Pro Ile Ser Arg Lys Glu Phe Val Arg Phe Leu Leu Ala Thr Pro Thr Lys Ser Glu Leu Arg Cys Gln Gly Leu Asn 215 Phe Ser Gly Ala Asp Leu Ser Arg Leu Asp Leu Arg Tyr Ile Asn Phe Lys Met Ala Asn Leu Ser Arg Cys Asn Leu Ala His Ala Asn Leu Cys Cys Ala Asn Leu Glu Arg Ala Asp Leu Ser Gly Ser Val Leu Asp Cys

270	
265 260 265 Ser Asn Ala Glu Gly Ala	
Ala Asn Leu Gln Gly Val Lys Met Leu Cys Ser Asn Ala Glu Gly Ala 285 275 280 275 Ser Leu Lys Leu Cys Asn Phe Glu Asp Pro Ser Gly Leu Lys Ala Asn 300 295 295 265 285 285 285 286 297 300 298 300 300 300 300 300	
Ser Leu Lys Leu Cys Asn Phe Glu Asp Pio Ser 300	
Ser Leu Lys Leu Cys Abn 295 300 290 295 Leu Glu Gly Ala Asn Leu Lys Gly Val Asp Met Glu Gly Ser Gln Met 320 315 310 315 Ala Lys Leu Lys	
Leu Glu Gly Ala Ash Leu Lys 315 310 315 315 317 318 318 319 319 319 310	
Leu Glu Gly Ala Ash 310 310 310 315 317 318 319 310 310 310 330 331 331 331	
325 Thr Leu Ala Gly Thr Asp Leu Glu Asn	
Asn Cys Asn Leu Arg Gly Ala Thr Leu Ala Gly Thr Asp Leu Glu Asn 325 Asn Cys Asn Leu Arg Gly Ala Thr Leu Ala Gly Thr Asp Leu Glu Asn 340 Cys Asp Leu Ser Gly Cys Asp Leu Gln Glu Ala Asn Leu Arg Gly Ser 360 360 370 380 380 380 380 380 380 380 380 380 38	
Cys Asp Leu Ser Gly Cys Asp Leu Gin	
Cys Asp Leu Ser Gly 535 360 365 355 Asn Val Lys Gly Ala Ile Phe Glu Glu Met Leu Thr Pro Leu His Met 375 380	
370 Ser Gln Ser Val Arg	
Ser Gin Ser Val 123	
<210> 28	
<211> 1246	
<212> DNA <213> H. sapiens	
(213) m. v-1	
<220>	
<221> CDS <222> (432)(1092)	
<223> K+Hnov44, splice 1	
	60
<400> 28 cagaaaacca cgcaggtcct tcttgatcat ctagaactga ccgctccgcc ttgccaggag cagaaaacca cgcaggtcct tcttgatcat ctagaactga ccgctccaggaa ggcggggggc tctgcagaac cacgtggcta gcctgcctga agttctcacc tctccaggaactac acttccgtgg tctgcagaac acgctgcg ctgggggctg ggggctcccg ctggggactcc acttccgtgg	120
cagaaaacca cgcaggteet gcctgcctga agttetcacc tetecaggaac ggggeteeg ettecaggactc acttecgtgg tetecaatgge tegcagetgge etgggggetg ggggeteeg etcagggtgaa agggagecat tetaatgge tegcagette tegcgccege aggggeatga etcagggtgaa agggagecat	180 240
tetgeagaac caegtyyeta setggggetg ggggeteeg etggggetea atgagetat tetaatgge tgeagetgeg etggggetga eteagtgaa aggggeatt atgetaage tetaacette ttgegeege aggggeatga eteagtgaa acaetteagg atgetaage etgggaaate acaetteagg	300
atgictaage titaggate atgeageet teageatee egigeaters agagagacag	360
tttctcagac ccctgaggg aggacagcct ttcctgcctc aggyattagagg gctggagagg	420 470
actacagtga tggagaccca ctagatgtgc acaagagggc ttc tca gtc cta atg	410
gcagccggag gcgctagggs 233 actacagaggct gccatctage getagaggct actacagtga tggagaccca ctagatgtgc acaagaggct tc tca gtc cta atg accgagccgt g atg ctg ggg ttt gcc atg atg ggc ttc tca gtc cta atg accgagccgt g atg ctg ggg ttt gcc atg atg ggc ttc tca gtc cta atg accgagccgt g atg ctg ggg ttt gcc atg atg ggc ttc tca gtc cta atg accgagccgt g atg ctg ggg ttt gcc atg atg ggc ttc tca gtc cta atg accgagccgt g atg ctg ggg ttt gcc atg atg ggc ttc tca gtc cta atg	
5	518
tto tto ttg ctc gga aca acc att cta aag cct ttt atg ctc agc att	210
tto tto ttg ctc gga aca acc att cta aag cct ttt atg ces as Ile Phe Phe Leu Leu Gly Thr Thr Ile Leu Lys Pro Phe Met Leu Ser Ile 20 25	
phe Phe Leu leu GI 7 20 25	566
atc atc dat	200
cag aga gaa tcg acc tgc act gcc atc cac aca gat acc acs Gln Arg Glu Glu Ser Thr Cys Thr Ala Ile His Thr Asp Ile Met Asp 45	
Gln Arg Glu Glu Ser 235 40	C1 A
gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac ggt cag gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc gac tgc ga	614
gac tgg ctg gac tgt gcc ttc acc tgt ggt gtg cac tgc cac gly Gln Asp Trp Leu Asp Cys Ala Phe Thr Cys Gly Val His Cys His Gly Gln 55 60	
Asp Trp Leu Asp Cys 55	
and cat cca ggt	662
ggg aag tac ccg tgt ctt cag gtg ttt ges Asn Leu Ser His Pro Gly	
Gly Lys Tyl Flo Clo To 70	
ong ata aat CCC	710
cag aaa gct ctc cta cat tat aat gaa gag gct gtc cag ata aat ccc	
Gln Lys Ala Leu leu 112 1 85	
aag tgc ttt tac aca cet aag tgc cac caa gat aga aat gat ttg ctc	758
too tac cac caa yar aya	
aag tgc ttt tac aca cct aag tgc out 48	

Lys Cys Phe Tyr Thr Pro Lys Cys His Gln Asp Arg Asn Asp Leu Leu 100 105	
aac agt gct ctg gac ata aaa gaa ttc ttc gat cac aaa aat gga act aac agt gct ctg gac ata aaa gaa ttc ttc gat cac aaa aat gga act aac agt gct ctg gac ata aaa gaa ttc ttc gat cac aaa aat gga act aac agt gct ctg gac ata aaa gaa ttc ttc gat cac aaa aat gga act aac agt gct ctg gac ata aaa gaa ttc ttc gat cac aaa aat gga act	806
ccc ttt tca tgc ttc tac agt cca gcc agc caa tct gaa gat gtc att pro Phe Ser Cys Phe Tyr Ser Pro Ala Ser Gln Ser Glu Asp Val Ile pro Phe Ser Cys Phe Tyr Ser Pro Ala Ser Gln Ser Glu Asp Val Ile	854
ctt ata aaa aag tat gac caa atg gct atc ttc cac tgt tta ttt tgg ctt ata aaa aag tat gac caa atg gct atc ttc cac tgt tta ttt tgg Leu Ile Lys Lys Tyr Asp Gln Met Ala Ile Phe His Cys Leu Phe Trp 150 150	902
cct tca ctg act ctg cta ggt ggt gcc ctg att gtt ggc atg gtg aga Pro Ser Leu Thr Leu Leu Gly Gly Ala Leu Ile Val Gly Met Val Arg 165	950
tta aca caa cac ctg tcc tta ctg tgt gaa aaa tat agc act gta gtc tta Thr Gln His Leu Ser Leu Leu Cys Glu Lys Tyr Ser Thr Val Val 180 185	998
aga gat gag gta ggt gga aaa gta cct tat ata gaa cag cat cag ttc Arg Asp Glu Val Gly Gly Lys Val Pro Tyr Ile Glu Gln His Gln Phe 195 200 205	1046
aaa ctg tgc att atg agg agg agc aaa gga aga gca gag aaa tct t aaa ctg tgc att atg agg agg agc aaa gga aga gca gag aaa tct t Lys Leu Cys Ile Met Arg Arg Ser Lys Gly Arg Ala Glu Lys Ser 220	1092
aagacggtgg ccaaattaaa gtgctggcct tcagatgtct gtgatttctg caactgagga cctaattatg cctgtctgca aactaataat gtaaaaggta ataattaaag tatcatattt tcatgtggga aaaaaaaaaa aaaaaaaaaa aaaa	1152 1212 1246
<210> 29 <211> 1111 <212> DNA <213> H. sapiens	
<220> <221> CDS <222> (297)(957) <223> K+Hnov44, splice 2	
<400> 29 aaaaaacatg acttgtggca ccagaagaga gccggggact tcaatccaag aaagcagagagagagacacaagagacaaa gaaggaccaa gcaaagaaga ctgtaccatg tcctaagctagagagagagagagagagagagagagagaga	ag 240
ctg ggg ttt gcc atg atg ggc ttc tca gtc cta atg ttc ttc ttg ctc Leu Gly Phe Ala Met Met Gly Phe Ser Val Leu Met Phe Phe Leu Leu 5	
gga aca acc att cta aag cct ttt atg ctc agc att cag aga gaa ga: Gly Thr Thr Ile Leu Lys Pro Phe Met Leu Ser Ile Gln Arg Glu Gl: 20 25 30	a 395 u

tog acc tgc act gcc atc cac aca gat atc atg gac gac tgg ctg gac 44 Ser Thr Cys Thr Ala Ile His Thr Asp Ile Met Asp Asp Trp Leu Asp 40 45	43
	91
	539
cta cat tat aat gaa gag gct gtc cag ata aat ccc aag tgc ttt tac cta cat tat aat gaa gag gct gtc cag ata aat ccc aag tgc ttt tac teu His Tyr Asn Glu Glu Ala Val Gln Ile Asn Pro Lys Cys Phe Tyr 90 95	587
aca cct aag tgc cac caa gat aga aat gat ttg ctc aac agt gct ctg Thr Pro Lys Cys His Gln Asp Arg Asn Asp Leu Leu Asn Ser Ala Leu 105 110	635
gac ata aaa gaa ttc ttc gat cac aaa aat gga act ccc ttt tca tgc Asp Ile Lys Glu Phe Phe Asp His Lys Asn Gly Thr Pro Phe Ser Cys 120 125	683
ttc tac agt cca gcc agc caa tct gaa gat gtc att ctt ata aaa aag ttc tac agt cca gcc agc caa tct gaa gat gtc att ctt ata aaa aag ttc tac agt cca gcc agc caa tct gaa gat gtc att ctt ata aaa aag ttc tac agt cca gcc agc caa tct gaa gat gtc att ctt ata aaa aag ttc tac agt cca gcc agc caa tct gaa gat gtc att ctt ata aaa aag 135	731
tat gac caa atg gct atc ttc cac tgt tta ttt tgg cct tca ctg act tat gac caa atg gct atc ttc cac tgt tta ttt tgg cct tca ctg act tat gac caa atg gct atc ttc cac tgt tta ttt tgg cct tca ctg act tat gac caa atg gct atc ttc cac tgt tta ttt tgg cct tca ctg act tat gac caa atg gct atc ttc cac tgt tta ttt tgg cct tca ctg act	779
ctg cta ggt ggt gcc ctg att gtt ggc atg gtg aga tta aca caa cac ctg cta ggt ggt gcc ctg att gtt ggc atg gtg aga tta aca caa cac Leu Cly Gly Ala Leu Ile Val Gly Met Val Arg Leu Thr Gln His 170 175	827
ctg tcc tta ctg tgt gaa aaa tat agc act gta gtc aga gat gag gta Leu Ser Leu Cys Glu Lys Tyr Ser Thr Val Val Arg Asp Glu Val 185 190	875
ggt gga aaa gta cct tat ata gaa cag cat cag ttc aaa ctg tgc att Gly Gly Lys Val Pro Tyr Ile Glu Gln His Gln Phe Lys Leu Cys Ile 200 205	923
atg agg agg agc aaa gga aga gca gag aaa tct t aagacggtgg atg agg agg agc aaa gga aga gca gag aaa tct t aagacggtgg Met Arg Arg Ser Lys Gly Arg Ala Glu Lys Ser 220	967
ccaaattaaa gtgctggcct tcagatgtct gtgatttctg caactgagga cctaattatg cctgtctgca aactaataat gtaaaaaggta ataattaaag tatcatattt tcatgtggga aaaaaaaaaa aaaaaaaaaa aaaa	1027 1087 1111
<210> 30 <211> 220 <212> PRT <213> H. sapiens	
<pre>Met Leu Gly Phe Ala Met Met Gly Phe Ser Val Leu Met Phe Phe Leu 15</pre>	

50

Leu Gly Thr Thr Ile Leu Lys Pro Phe Met Leu Ser Ile Gln Arg Glu 30 25 30 777 Leu	
Leu Gly Thr Thr Ile Leu Lys 25	
20 25 Glu Ser Thr Cys Thr Ala Ile His Thr Asp Ile Met Asp Asp Trp Leu 45 40 40 Cly Lys TVI	
35 Asp Cys Ala Phe Thr Cys Gly Val His Cys His Gly Gln Gly Lys Tyr 60	
55 50 Fro Cys Leu Gln Val Phe Val Asn Leu Ser His Pro Gly Gln Lys Ala Pro Cys Leu Gln Val Phe Val Asn Leu Ser His Pro Gly Gln Lys Ala 80 75 80 60 60 60 60 60 60 60 60 60 60 60 60 60	
Pro Cys Leu Gin vai Pile vai - 75	
70 75 76 76 76 76 77 77 78 78 79 79 79 79 79 79 79 79 79 79 79 79 79	
Leu Leu His 191 Asi Ser Ala	
85 Tyr Thr Pro Lys Cys His Gln Asp Arg Asn Asp Leu Leu Asn Ser Ala 110 105 105 107 108 109 109 100 100 100 100 100 100 100 100	
100 His Lys Asn Gly Thr Pro Phe Ser	
105 100 Leu Asp Ile Lys Glu Phe Phe Asp His Lys Asn Gly Thr Pro Phe Ser 125 120 125 127 128 129 129 120 125	
115 Cys Phe Tyr Ser Pro Ala Ser Gln Ser Glu Asp Val Ile Leu Ile Lys 135 140 135 140 140	
Cys Phe Tyr Ser Flo All 135	
135 130 130 135 130 135 140 140 140 150 160 150 150 150 150 150 150 150 150 150 15	
150 150 Arg Leu Thr Gln	
150 155 145 150 150 175 Thr Leu Leu Gly Gly Ala Leu Ile Val Gly Met Val Arg Leu Thr Gln 175 170 175 170 175 175	
165 Tyr Ser Thr Val Val Arg Asp Glu	
Thr Leu Bed 617 170 165 His Leu Ser Leu Leu Cys Glu Lys Tyr Ser Thr Val Val Arg Asp Glu 190 185	
Val Gly Gly Lys Val Pro Tyr Ile Glu Gln His Gln Phe Lys Leu Cys 205	
Val Gly Gly Lys Val 413 17 200 205	
195 Ile Met Arg Arg Ser Lys Gly Arg Ala Glu Lys Ser 215 220	
215 210	
•••	
<210> 31	
<211> 22	
<212> DNA <213> Artificial Sequence	
<213> AFTILICIAL SOLUTION	
<220>	
<223> consensus sequences	
	22
<400> 31	
tatecacate aatggacaaa ge	
<210> 32	
<211> 20	
-212 NAA	
<213> Artificial Sequence	
22	20
<400> 32 tgcataactg gctgggtgta	
tgcataacty googss-s	
<210> 33	
<211> 22	
<212> DNA	
<213> Artificial Sequence	•
<400> 33	22
tgacatcact ggatgaactt ga	
tyacatours 55 F	
<210> 34	
<211> 20	
<212> DNA <213> Artificial Sequence	
<213> ALUILIUM	.00
<400> 34	20
rgcctgcaaa gtttgaacat	
51	

<210> 35	
<211> 22	
<212> DNA <213> Artificial Sequence	
<213> Artificial Sequence	
<400> 35	22
tgacatcact ggatgaactt ga	22
tgacaccacc 55-55-55	
<210> 36	
<211> 20	
<212> DNA	
<213> Artificial Sequence	
100. 26	20
<pre><400> 36 tgcctgcaaa gtttgaacat</pre>	20
Egectgeada geetgaavas	
<210> 37	
<211> 20	
<212> DNA	
<213> Artificial Sequence	
100: 37	20
<400> 37 acctggtggt atggaagcat	20
acctggtggt atggaageas	
<210> 38	
<211> 19	
<212> DNA	
<213> Artificial Sequence	
20	
<400> 38 tttctcctgg cctctaccc	19
Effeteetigg effetiation	
<210> 39	
<211> 19	
c212> DNA	
<213> Artificial Sequence	
<400> 39	19
tccctcttgg gtgaccttc	19
Eddarding Angusta	
<210> 40	
<211> 20	
<212> DNA	
<213> Artificial Sequence	
<400> 40	20
atctttgtca gccaccagct	20
accepted goodenson	
<210> 41	
<211> 24	
<212> DNA	
<213> Artificial Sequence	
<400> 41	24
aggtgtgctg ccatctgctg ttcg	24
aggracaca comments in	
<210> 42	
<211> 24	
<212> DNA	
213> Artificial Sequence	

	24
<400> 42 agcctatcct ctctgagagt cagg	
<210> 43	
<211> 21	•
THE THIRD	
<212> DNA <213> Artificial Sequence	
	21
<400> 43	
aagcagagta ctcatgatgc c	
<210> 44	
<211> 20	
<212> DNA	
<212> DNA <213> Artificial Sequence	
<400> 44	20
tctggtagac agtacagtgg	
<210> 45	
<211> 20	
<212> DNA	
<212> DAR <213> Artificial Sequence	20
<400> 45	20
catttggctg gtccaagatg	
<210> 46	
<211> 20	
and. DNA	
<213> Artificial Sequence	
<213> ALCIE 2011	20
<400> 46	
agtcattggt agggaggtac	
<210> 47	
<211> 20	
<212> DNA	
<212> DMA <213> Artificial Sequence	20
<400> 47	20
catgetteta cagtecagee	
<210> 48	
<211> 20	•
<212> DNA <213> Artificial Sequence	
<213> ALLII10200	20
<400> 48	
ggtcctcagt tgcagaaatc	
<210> 49	
<211> 45	
<212> DNA Ambificial Sequence	
<212> bha <213> Artificial Sequence	
<400> 49	45
<400> 49 tggtgggctg tggtgaccat gacaactgtg ggctatgggg acatg	
<210> 50	

<211> 45	
-2125 DNA	
<213> Artificial Sequence	
<400> 50	45
<400> 50 tggtgggcag tggtcaccat gaccactgtg ggctacgggg acatg	
<210> 51 <211> 45	
<212> DNA	
<213> Artificial Sequence	
<400> 51	45
<400> 51 tggtgggcag tcgtctccat gacaactgta ggctatggag acatg	
<210> 52	
<211> 45	
<212> DNA <213> Artificial Sequence	
<213> Artificial Sequence	
<400> 52	45
<pre><400> 52 tggtgggcag tggtaaccat gacaacagtg ggttacggcg atatg</pre>	
ragragame assessment of the control	
<210> 53	
<211> 45	
<212> DNA	
<213> Artificial Sequence	
<400> 53	45
<400> 53 tggtgggetg tggtcaccat gacgaccetg ggctatggag acatg	
tggtgggccg cgg .	
<210> 54	
<211> 45	
<212> DNA	
<213> Artificial Sequence	
<400> 54	45
<400> 54 tggtgggggg tggtcacagt caccaccatc ggctatgggg acaag	
C33-533333 20	
<210> 55	
<211> 45	
<212> DNA <213> Artificial Sequence	
(213) ALUILIUGA BUJANTA	
<400> 55	45
<400> 55 tggtgggcag tggtcaccat gaccacggtt ggctatgggg acatg	
<210> 56	
<211> 45	
<212> DNA <213> Artificial Sequence	
(ST2) MILLITOTOT CONTINUE	
<400> 56	45
<pre><400> 56 tggtgggccg tggtcaccat gacgaccctg ggctatggag acatg</pre>	
<210> 57	
<211> 45	
<212> DNA <213> Artificial Sequence	
<213> AITHIGHAL SOQUENOS	
<400> 57	

•	45
tggtgggctg tggtcaccat gacgacactg ggctacggag acatg	
<210> 58	
<211> 45	
<212> DNA	
<213> Artificial Sequence	
<400> 58	45
<400> 58 tggtgggetg tggtgaccat gacaactgtg ggctatgggg acatg	
<210> 59	
<211> 47	
-212> DNA	
<213> Artificial Sequence	
<400> 59	47
<400> 59 ttcctgttct ccattgagac cgaaacaacc attgggtatg gcttccg	
<210> 60	
<211> 47	
<212> DNA	
<213> Artificial Sequence	
<400> 60	47
<400> 60 tttttattct caatagagac agaaaccacc attggttatg gctaccg	
<210> 61	
<211> 47	
212 DNA	
<213> Artificial Sequence	
(213) RICII-	47
<400> 61 areassaacc ataggetatg gtttcag	**
<pre><400> 61 ttcctcttct ccattgagac ccagacaacc ataggctatg gtttcag</pre>	
<210> 62	
<211> 47	
-212 DNA	
<213> Artificial Sequence	
	47
<400> 62 ttcctgttct cggtggagac gcagacgacc atcggctatg ggttccg	
FEGGEGEGE GAARANA T	
<210> 63	
<211> 47	
<212> DNA	
<213> Artificial Sequence	
<400> 63	47
<400> 63 ttcctcttct cccttgaatc ccaaaccacc attggctatg gcttccg	
<210> 64	
<211> 47	
-212 DNA	
<213> Artificial Sequence	
•	47
<400> 64	47
<pre><400> 64 tttctctttt ccctggaatc ccagacaacc attggctatg gagtccg</pre>	
<210> 65	
<211> 47	
<212> DNA	

```
<213> Artificial Sequence
                                                                       47
ttccttttct ccattgaggt ccaagtgact attggctttg gggggcg
      <210> 66
      <211> 47
      <212> DNA
      <213> Artificial Sequence
                                                                        47
tttctcttct ccattgaagt tcaagttacc attgggtttg gagggag
      <210> 67
      <211> 50
       <212> DNA
       <213> Artificial Sequence
                                                                        50
 gegetetact teacetteag eagecteace agtgtggget teggeaacgt
       <210> 68
       <211> 15
       <212> PRT
       <213> Artificial Sequence
       <220>
       <223> consensus sequences
  Trp Trp Ala Val Val Ser Met Thr Thr Val Gly Tyr Gly Asp Met
                  5
        <210> 69
        <211> 15
        <212> PRT
        <213> Artificial Sequence
   Trp Trp Ala Val Val Thr Met Thr Thr Leu Gly Tyr Gly Asp Met
    1
         <210> 70
         <211> 15
         <212> PRT
         <213> Artificial Sequence
    Trp Trp Gly Val Val Thr Val Thr Thr Ile Gly Tyr Gly Asp Lys
                    5
          <210> 71
          <211> 15
          <212> PRT
          <213> Artificial Sequence
    Trp Trp Ala Val Val Thr Met Thr Thr Val Gly Tyr Gly Asp Met
                     5
```

56

<210> 72

```
<211> 15
     <212> PRT
     <213> Artificial Sequence
Phe Leu Phe Ser Ile Glu Val Gln Val Thr Ile Gly Phe Gly Gly
1
     <210> 73
      <211> 15
      <212> PRT
      <213> Artificial Sequence
Phe Leu Phe Ser Leu Glu Ser Gln Thr Thr Ile Gly Tyr Gly Val
                 5
 1
      <210> 74
       <211> 15
       <212> PRT
       <213> Artificial Sequence
 Phe Leu Phe Ser Ile Glu Thr Glu Thr Thr Ile Gly Tyr Gly Tyr
  1
       <210> 75
       <211> 15
        <212> PRT
        <213> Artificial Sequence
  Phe Leu Phe Ser Ile Glu Thr Gln Thr Thr Ile Gly Tyr Gly Phe
                   5
   1
        <210> 76
        <211> 15
         <212> PRT
         <213> Artificial Sequence
   Phe Leu Phe Ser Val Glu Thr Gln Thr Thr Ile Gly Tyr Gly Phe
         <210> 77
         <211> 15
         <212> PRT
         <213> Artificial Sequence
    Phe Leu Phe Ser Leu Glu Ser Gln Thr Thr Ile Gly Tyr Gly Phe
     1
          <210> 78
          <211> 15
          <212> PRT
          <213> Artificial Sequence
     Phe Leu Phe Ser Ile Glu Thr Glu Thr Thr Ile Gly Tyr Gly Phe
      1
                                         57
```

<210> 79 <211> 16	
<212> PRT <213> Artificial Sequence	
<pre><400> 79 Ala Leu Tyr Phe Thr Phe Ser Ser Leu Thr Ser Val Gly Phe Gly Asn 1</pre>	٠
<210> 80 <211> 2571 <212> DNA <213> H. sapiens	
<220> <221> CDS <222> (110) (1285)	
<pre><400> 80 gctgccgcgc ctgtagcact cccggaactg gaactaggtg ccagacggtc cggaggcggg ggccacgtca gcggggccac ccagggctcg cggggtcccg gtgggtgcc atg cgg agg</pre>	60 118
ggc gcg ctt ctg gcg ggc gcc ttg gcc gcg tac gcc gcg tac ctg gtg Gly Ala Leu Leu Ala Gly Ala Leu Ala Ala Tyr Ala Ala Tyr Leu Val 10	166
ctg ggc gcg ctg ttg gtg gcg cgg ctg gag ggg ccg cac gaa gcc agg ctg ggc gcg ctg ttg gtg gcg cgg ctg gag ggg ccg cac gaa gcc agg Leu Gly Ala Leu Leu Val Ala Arg Leu Glu Gly Pro His Glu Ala Arg Leu Gly Ala Leu Leu Val Ala Arg Leu Glu Gly Pro His Glu Ala Arg 25 30	214
ctc cga gcc gag ctg gag acg ctg cgg gcg cag ctg ctt cag cgc agc ctc cga gcc gag ctg gag acg ctg cgg gcg cag ctg ctt cag cgc agc Leu Arg Ala Glu Leu Glu Thr Leu Arg Ala Gln Leu Leu Gln Arg Ser 40 45 50	262
ccg tgt gtg gct gcc ccc gcc ctg gac gcc ttc gtg gag cga gtg ctg Pro Cys Val Ala Ala Pro Ala Leu Asp Ala Phe Val Glu Arg Val Leu 55 60 65	310
gcg gcc gga cgg ctg ggg cgg gtc gtg ctt gct aac gct tcg ggg tcc Ala Ala Gly Arg Leu Gly Arg Val Val Leu Ala Asn Ala Ser Gly Ser 70 75 80	358
gcc aac gcc tcg gac ccc gcc tgg gac ttc gcc tct gct ctc ttc ttc Ala Asn Ala Ser Asp Pro Ala Trp Asp Phe Ala Ser Ala Leu Phe Phe	406
gcc agc acg ctg atc acc acc gtg ggc tat ggg tac aca acg cca ctg Ala Ser Thr Leu Ile Thr Thr Val Gly Tyr Gly Tyr Thr Thr Pro Leu 105 110 115	454
act gat gcg ggc aag gcc ttc tcc atc gcc ttt gcg ctc ctg ggc gtg Thr Asp Ala Gly Lys Ala Phe Ser Ile Ala Phe Ala Leu Leu Gly Val 120 125 130	502
ccg acc acc atg ctg ctg ctg acc gcc tca gcc cag cgc ctg tca ctg Pro Thr Thr Met Leu Leu Thr Ala Ser Ala Gln Arg Leu Ser Leu 135	550

WO 99/43696 PCT/US99/03826

ctg ctg act cac gtg ccc ctg tct tgg ctg agc atg cgt tgg ggc tgg Leu Leu Thr His Val Pro Leu Ser Trp Leu Ser Met Arg Trp Gly Trp 150 155	598
gac ccc cgg cgg gcg gcc tgc tgg cac ttg gtg gcc ctg ttg ggg gtc Asp Pro Arg Arg Ala Ala Cys Trp His Leu Val Ala Leu Leu Gly Val 165 170 175	646 .
gta gtg acc gtc tgc ttt ctg gtg ccg gct gtg atc ttt gcc cac ctc Val Val Thr Val Cys Phe Leu Val Pro Ala Val Ile Phe Ala His Leu 185 190 195	694
gag gag gcc tgg agc ttc ttg gat gcc ttc tac ttc tgc ttt atc tct Glu Glu Ala Trp Ser Phe Leu Asp Ala Phe Tyr Phe Cys Phe Ile Ser 200 205 210	742
ctg tcc acc atc ggc ctg ggc gac tac gtg ccc ggg gag gcc cct ggc Leu Ser Thr Ile Gly Leu Gly Asp Tyr Val Pro Gly Glu Ala Pro Gly	790
cag ccc tac cgg gcc ctc tac aag gtg ctg gtc aca gtc tac ctc ttc Gln Pro Tyr Arg Ala Leu Tyr Lys Val Leu Val Thr Val Tyr Leu Phe	838
ctg ggc ctg gtg gcc atg gtg ctg gtg ctg cag acc ttc cgc cac gtg ctg ggc ctg gtg gcc atg gtg ctg ctg cag acc ttc cgc cac gtg Leu Gly Leu Val Ala Met Val Leu Val Leu Gln Thr Phe Arg His Val 250 255	886
tcc gac ctc cac ggc ctc acg gag ctc atc ctg ctg ccc cct ccg tgc_ Ser Asp Leu His Gly Leu Thr Glu Leu Ile Leu Leu Pro Pro Cys 275	934
cct gcc agt ttc aat gcg gat gag gac gat cgg gtg gac atc ctg ggc Pro Ala Ser Phe Asn Ala Asp Glu Asp Asp Arg Val Asp Ile Leu Gly 280 285	982
ccc cag ccg gag tcg cac cag caa ctc tct gcc agc tcc cac acc gac Pro Gln Pro Glu Ser His Gln Gln Leu Ser Ala Ser Ser His Thr Asp 295 300 305	1030
tac gct tcc atc ccc agg tag ctg ggg cag cct ctg cca ggc ttg ggt Tyr Ala Ser Ile Pro Arg * Leu Gly Gln Pro Leu Pro Gly Leu Gly	1078
gtg cct ggc ctg gga ctg agg ggt cca ggc gac cag agc tgg ctg tac Val Pro Gly Leu Gly Leu Arg Gly Pro Gly Asp Gln Ser Trp Leu Tyr 325 330 335	1126
agg aat gtc cac gag cac agc agg tga tct tga ggc ctt gcc gtc cac Arg Asn Val His Glu His Ser Arg * Ser * Gly Leu Ala Val His 340 345	1174
cgt ctc tcc ttt gtt tcc cag cat ctg gct ggg atg tga agg gca gca Arg Leu Ser Phe Val Ser Gln His Leu Ala Gly Met * Arg Ala Ala 355 360 365	1222
ctc cct gtc ccc atg tcc cgg gct cca ctg ggc acc aac ata acc ttg Leu Pro Val Pro Met Ser Arg Ala Pro Leu Gly Thr Asn Ile Thr Leu 370 375 380	1270
tte tet gte ett tet etcateetet ttacaetgtg tetetetgge tetetggeat	1325

Phe Ser Val Leu Ser 385

totogotgoo totgtottto cotottgotg tototgttto toattotott toatgttoog tetgtgtete teaattaace actegteaac tgetgattet actgggetgt gggeteagae ctcatttcag gcaccagatt ggtcgctaca ccctggacaa gtgactgccc gtctctgagc 1505 cttgatttcc tcagctgcca aatgggaaga atagaagaat ttgcccctaa acccctcctg 1565 tgtgctggcc ctgtgctaga cagtgctgga gacatagttg ggggtggaga actgccctta 1625 tggagettge agtecagtga ggtggacaga cetgtececa gacagtgatg geecaaaatg gtcaggactt taatggagga ggtgaggtgt tgaaagcaca ggcagagtgg tcagggctga 1745 agtcggagaa gcatagggac taggcccaat ccagcctgga aagtcaggga ggacttccta 1805 gaggaaggga catcgaacta agacctgaac tatgagaaat aggcaggaag aagttgtacc 1865 1925 tgactcattt ttetcaggtg tetecaggga geaggaceca tggagggace cetggtgtag gcctgggcga tagactette etcagcagee tggcaggcag gaaacagaca taggaceeca geccagatet gaatggeatg ggaggtgetg ceettaacca tgacaccatt gtaagagetg 2045 tecacatttg tatgttgtge cetggaatca geetggttga geteaaatce caaettagee 2105 acgtctggcc tgtgtccttg ggcagtcaca ctacctctct gattttgttt ccttatctgt 2165 aaaatggtga tcatcataat acaacttcaa aaggatttca ggctgagtgt ggtggctcac 2225 gcctatacac ccagcacttt ggaaggctga ggaaggagga tcgcttgagg ccaggagttt 2285 gagactagee taggeaacae agtgaggeet tateteaaca acaaccacaa aatetaaaaa 2345 ttagctgggt gtggtggtgc atgcctgtga tcctggctac ttcagaggct gaggtggaag 2405 gatcacttga ggccaggagt ttgaggctgc agtgagttat gatggcactg ctgcactcca 2465 geetgeggga cagagtgaga eeetgtetga aagaaagaga gaaagaaaga aagaaagaga 2525 2571

<210> 81

<211> 388

<212> PRT

<213> H. sapiens

Met Arg Arg Gly Ala Leu Leu Ala Gly Ala Leu Ala Ala Tyr Ala Ala Tyr Leu Val Leu Gly Ala Leu Leu Val Ala Arg Leu Glu Gly Pro His Glu Ala Arg Leu Arg Ala Glu Leu Glu Thr Leu Arg Ala Gln Leu Leu Gln Arg Ser Pro Cys Val Ala Ala Pro Ala Leu Asp Ala Phe Val Glu Arg Val Leu Ala Ala Gly Arg Leu Gly Arg Val Val Leu Ala Asn Ala Ser Gly Ser Ala Asn Ala Ser Asp Pro Ala Trp Asp Phe Ala Ser Ala Leu Phe Phe Ala Ser Thr Leu Ile Thr Thr Val Gly Tyr Gly Tyr Thr Thr Pro Leu Thr Asp Ala Gly Lys Ala Phe Ser Ile Ala Phe Ala Leu 120 Leu Gly Val Pro Thr Thr Met Leu Leu Leu Thr Ala Ser Ala Gln Arg Leu Ser Leu Leu Thr His Val Pro Leu Ser Trp Leu Ser Met Arg Trp Gly Trp Asp Pro Arg Arg Ala Ala Cys Trp His Leu Val Ala Leu 165 170 175 Leu Gly Val Val Val Thr Val Cys Phe Leu Val Pro Ala Val Ile Phe 185 Ala His Leu Glu Glu Ala Trp Ser Phe Leu Asp Ala Phe Tyr Phe Cys

200 Phe Ile Ser Leu Ser Thr Ile Gly Leu Gly Asp Tyr Val Pro Gly Glu Ala Pro Gly Gln Pro Tyr Arg Ala Leu Tyr Lys Val Leu Val Thr Val

215

230

The Val Ala Met Val Leu Val Leu Gin The File	
Tyr Leu Phe Leu Gly Leu Val Ala Met Val Leu Val Leu Gln Thr Phe 255 245 Arg His Val Ser Asp Leu His Gly Leu Thr Glu Leu Ile Leu Leu Pro 265 265 267 268 269 270 280 280 280 280 280 280 280 280 280 28	
Arg His Val Ser Asp Leu His Gly 265 260 260 260 Asp Asp Arg Val Asp	
Pro Pro Cys Pro Ala Ser Phe Asn Ala Asp Glu Asp Asp Arg Val Asp 275 11e Leu Gly Pro Gln Pro Glu Ser His Gln Gln Leu Ser Ala Ser Ser 295 207 208 209 209 209 200 200 200 200	,
The Leu Gly Pro Gin Pro Giv 295 290 His Thr Asp Tyr Ala Ser Ile Pro Arg Leu Gly Gln Pro Leu Pro Gly 320 310 310 310 310 300 320 320	
His Thr Asp Tyr Ala Ser 11e Pro Ala 315 310 310 310 310 320	
His Thr Asp 191 All 310 310 305 Leu Gly Val Pro Gly Leu Gly Leu Arg Gly Pro Gly Asp Gln Ser Trp 335 Leu Gly Val Pro Gly Leu Ala Val His	
Leu Gly Val Plo Gly Leu 330 325 Leu Tyr Arg Asn Val His Glu His Ser Arg Ser Gly Leu Ala Val His 350 345 340 345 340 340 340 340 340 341 340 340 341 340 340 340 340 340 340	
Leu Tyr Arg Ash Vol 345 340 Arg Leu Ser Phe Val Ser Gln His Leu Ala Gly Met Arg Ala Ala Leu 365 360 361 365 360 361 361 361 365	
Arg Leu Ser Phe Val 360 355 355 Pro Val Pro Met Ser Arg Ala Pro Leu Gly Thr Asn Ile Thr Leu Phe 370 370 380	
Ser Val Leu Ser 385	
<210> 82 <211> 3300	•
<2.12> DNA	
<213> H. sapiens	
<220> <221> CDS	
<222> (50)(1285)	
<pre><400> 82 aaatgcctgc ccgtgcagct cggagcgcgc agcccgtctc tgaataaga atg gcg gca Met Ala Ala 1</pre>	58
1	
PPS PDD SEE pad a	106
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 15 10 15	106 154
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 10 15 ctc tcg ttt tcc acg aaa ccc aca gtg ctt gct tcc cgg gtg gag agt Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg Val Glu Ser	
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 10 15 ctc tcg ttt tcc acg aaa ccc aca gtg ctt gct tcc cgg gtg gag agt Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg Val Glu Ser 25 20 20 25 26 27 28 28 28 28 28 28 28 28 28	
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 15 ctc tcg ttt tcc acg aaa ccc aca gtg ctt gct tcc cgg gtg gag agt Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg Val Glu Ser 20 25 gac acg acc att aat gtt atg aaa tgg aag acg gtc tcc acg ata ttc Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe	154
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 10 ctc tcg ttt tcc acg aaa ccc aca gtg ctt gct tcc cgg gtg gag agt Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg Val Glu Ser 25 gac acg acc att aat gtt atg aaa tgg aag acg gtc tcc acg ata ttc Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe 45	154
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 10 ctc tcg ttt tcc acg aaa ccc aca gtg ctt gct tcc cgg gtg gag agt Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg Val Glu Ser 25 gac acg acc att aat gtt atg aaa tgg aag acg gtc tcc acg ata ttc Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe 45	15 4 202
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 10 15 ctc tcg ttt tcc acg aaa ccc aca gtg ctt gct tcc cgg gtg gag agt Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg Val Glu Ser 25 gac acg acc att aat gtt atg aaa tgg aag acg gtc tcc acg ata ttc Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe 40 ctg gtg gtt gtc ctc tat ctg atc atc gga gcc acc gtg ttc aaa gca Leu Val Val Val Leu Tyr Leu Ile Ile Gly Ala Thr Val Phe Lys Ala 60 65	15 4 202
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 10 15 ctc tcg ttt tcc acg aaa ccc aca gtg ctt gct tcc cgg gtg gag agt Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg Val Glu Ser 25 gac acg acc att aat gtt atg aaa tgg aag acg gtc tcc acg ata ttc Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe 40 ctg gtg gtt gtc ctc tat ctg atc atc gga gcc acc gtg ttc aaa gca Leu Val Val Val Leu Tyr Leu Ile Ile Gly Ala Thr Val Phe Lys Ala 60 65	154 202 250
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 10 15 ctc tcg ttt tcc acg aaa ccc aca gtg ctt gct tcc cgg gtg gag agt Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg Val Glu Ser 25 gac acg acc att aat gtt atg aaa tgg aag acg gtc tcc acg ata ttc Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe 45 ctg gtg gtt gtc ctc tat ctg atc atc gga gcc acc gtg ttc aaa gca Leu Val Val Val Leu Tyr Leu Ile Ile Gly Ala Thr Val Phe Lys Ala ttg gag cag cct cat gag att tca cag agg acc acc att gtg atc cag Leu Glu Gln Pro His Glu Ile Ser Gln Arg Thr Thr Ile Val Ile Gln 75	154 202 250
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 10 15 ctc tcg ttt tcc acg aaa ccc aca gtg ctt gct tcc cgg gtg gag agt Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg Val Glu Ser 25 gac acg acc att aat gtt atg aaa tgg aag acg gtc tcc acg ata ttc Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe 45 ctg gtg gtt gtc ctc tat ctg atc atc gga gcc acc gtg ttc aaa gca Leu Val Val Val Leu Tyr Leu Ile Ile Gly Ala Thr Val Phe Lys Ala ttg gag cag cct cat gag att tca cag agg acc acc att gtg atc cag Leu Glu Gln Pro His Glu Ile Ser Gln Arg Thr Thr Ile Val Ile Gln 75	154 202 250 298
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 10 15 ctc tcg ttt tcc acg aaa ccc aca gtg ctt gct tcc cgg gtg gag agt Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg Val Glu Ser 20 25 30 35 gac acg acc att aat gtt atg aaa tgg aag acg gtc tcc acg ata ttc Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe 40 45 50 ctg gtg gtt gtc ctc tat ctg atc atc gga gcc acc gtg ttc aaa gca Leu Val Val Leu Tyr Leu Ile Ile Gly Ala Thr Val Phe Lys Ala 55 60 65 ttg gag cag cct cat gag att tca cag agg acc acc act att gtg atc cag Leu Glu Gln Pro His Glu Ile Ser Gln Arg Thr Thr Ile Val Ile Gln 70 75 80 aag caa aca ttc ata tcc caa cat tcc tgt gtc aat tcg acg gag ctg Lys Gln Thr Phe Ile Ser Gln His Ser Cys Val Asn Ser Thr Glu Leu	154 202 250 298
cct gac ttg ctg gat cct aaa tct gcc gct cag aac tcc aaa ccg agg Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser Lys Pro Arg 10 15 ctc tcg ttt tcc acg aaa ccc aca gtg ctt gct tcc cgg gtg gag agt Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg Val Glu Ser 25 gac acg acc att aat gtt atg aaa tgg aag acg gtc tcc acg ata ttc Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser Thr Ile Phe 40 ctg gtg gtt gtc ctc tat ctg atc atc gga gcc acc gtg ttc aaa gca Leu Val Val Leu Tyr Leu Ile Ile Gly Ala Thr Val Phe Lys Ala 55 ttg gag cag cct cat gag att tca cag agg acc acc att gtg atc cag Leu Glu Gln Pro His Glu Ile Ser Gln Arg Thr Thr Ile Val Ile Gln 75 aag caa aca ttc ata tcc caa cat tcc tgt gtc aat tcg acg gag ctg Lys Gln Thr Phe Ile Ser Gln His Ser Cys Val Asn Ser Thr Glu Leu	154 202 250 298

61

	105	110	115
Pro Leu Gly Asn	and the aat cas	a atc agt cac tgg gat 1 Ile Ser His Trp Asp 125	ttg gga agt 442 Leu Gly Ser 130
	ago act ati	t att aca acc ata gga l Ile Thr Thr Ile Gly 140	ttt gga aac 490 Phe Gly Asn 145
atc tca cca cgc Ile Ser Pro Arg 150	aca gaa ggc gg Thr Glu Gly Gl 15	c aaa ata ttc tgt atc y Lys Ile Phe Cys Ile 5 160	atc tat gcc 538 Ile Tyr Ala
tta ctg gga att Leu Leu Gly Ile 165	ccc ctc ttt gg Pro Leu Phe Gl 170	t ttt ctc ttg gct gga y Phe Leu Leu Ala Gly 175	gtt gga gat 586 Val Gly Asp
cag cta ggc acc Gln Leu Gly Thr 180	ata ttt gga aa Ile Phe Gly Ly 185	a gga att gcc aaa gtg s Gly Ile Ala Lys Val 190	gaa gat acg 634 Glu Asp Thr 195
ttt att aag tgg Phe Ile Lys Trp	aat gtt agt ca Asn Val Ser Gl 200	ng acc aag att cgc atc In Thr Lys Ile Arg Ile 205	e atc tca aca 682 e Ile Ser Thr 210
atc ata ttt ata Ile Ile Phe Ile 215	Leu Phe Gly C	gt gta ctc ttt gtg gct ys Val Leu Phe Val Ala 220	ctg cct gcg 730 A Leu Pro Ala 225
atc ata ttc aaa Ile Ile Phe Lys 230	His Ile Glu G	gc tgg agt gcc ctg ga ly Trp Ser Ala Leu As 35 24	
ttt gtg gtt ato Phe Val Val Ile 245	c act cta aca a Thr Leu Thr T 250	ct att gga ttt ggt ga hr Ile Gly Phe Gly As 255	c tac gtt gca 826 p Tyr Val Ala
ggt gga tcc gat Gly Gly Ser As 260	t att gaa tat o p Ile Glu Tyr I 265	etg gac ttc tat aag co eu Asp Phe Tyr Lys Pr 270	t gtc gtg tgg 874 o Val Val Trp 275
ttc tgg atc ct Phe Trp Ile Le	t gta ggg ctt g u Val Gly Leu A 280	gct tac ttt gct gct gt Ala Tyr Phe Ala Ala Va 285	c ctg agc atg 922 11 Leu Ser Met 290
att gga gat tg Ile Gly Asp Tr 29	p Leu Arg Val .	ata tct aaa aag aca aa Ile Ser Lys Lys Thr Ly 300	aa gaa gag gtg 970 ys Glu Glu Val 305
gga gag ttc ag Gly Glu Phe Ar 310	g Ala His Ala .	gct gag tgg aca gcc a Ala Glu Trp Thr Ala A 315 3	ac gtc aca gcc 1018 gn Val Thr Ala 20
gaa ttc aaa ga Glu Phe Lys Gl 325	aa acc agg agg lu Thr Arg Arg 330	cga ctg agt gtg gag a Arg Leu Ser Val Glu I 335	tt tat gac aag 1066 le Tyr Asp Lys
ttc cag cgg g Phe Gln Arg A 340	cc acc tcc atc la Thr Ser Ile 345	aag egg aag ete teg g Lys Arg Lys Leu Ser A 350	ca gaa ctg gct 1114 la Glu Leu Ala 355

gga aac cac aat cag gag ctg act cct tgt agg agg acc ctg tca gtg Gly Asn His Asn Gln Glu Leu Thr Pro Cys Arg Arg Thr Leu Ser Val 365 370	
aac cac ctg acc agc gag agg gat gtc ttg cct ccc tta ctg aag act 1210 Asn His Leu Thr Ser Glu Arg Asp Val Leu Pro Pro Leu Leu Lys Thr 375 380 385	
gag agt atc tat ctg aat ggt ttg acg cca cac tgt gct ggt gaa gag 1258 Glu Ser Ile Tyr Leu Asn Gly Leu Thr Pro His Cys Ala Gly Glu Glu 395 400	
390	
att gct gtg att gag aac atc aaa tag ccctctcttt aaataacctt 1305 Ile Ala Val Ile Glu Asn Ile Lys * 405 410	
aggcatagcc ataggtgagg acttetetat getetttatg actgttgetg gtagcatetat 1425 aggcatagcc ataggtgagg acttetetat getetttatg actgttgetg gtagcatetat 1425 1485	
aaataatttc cctaaatata attgcaattcag attagggtct tgaaaaataa acccagaatc 2005	
	j
	;
aaattttag aaagtcaggc tcttttagaa agaaagctac acccatttee double 3225 gttccgaaaa tttatatggt ggaatgcgcc atgtataaac tgtgaattgt attgacaaat 3285 gttccgaaaa tttatatggt aaaaaaaaaaa aaaaaaaaaa	
gttccgaaaa tttatatggt ggaatgcgcc atgtataaac tgtgaaceg aaaaaaaaaa 3289 aaagtttgta attaaagtca aaaaaaaaaa aaaaaaaaaa)
aaaaaaaaa aaaaa	
aaadaaaaa	•
<210> 83	
<211> 411	
<212> PRT	
<213> H. sapiens	
<pre><400> 83 Met Ala Ala Pro Asp Leu Leu Asp Pro Lys Ser Ala Ala Gln Asn Ser</pre>	
Met Ala Ala Plo App 10 10 10 In Italy Ala Ser Arg	
10 1 5 1 Lys Pro Arg Leu Ser Phe Ser Thr Lys Pro Thr Val Leu Ala Ser Arg	
Lys P10 A2 200 63	

```
25
Val Glu Ser Asp Thr Thr Ile Asn Val Met Lys Trp Lys Thr Val Ser
                40
Thr Ile Phe Leu Val Val Leu Tyr Leu Ile Ile Gly Ala Thr Val
  50 55
Phe Lys Ala Leu Glu Gln Pro His Glu Ile Ser Gln Arg Thr Thr Ile
Val Ile Gln Lys Gln Thr Phe Ile Ser Gln His Ser Cys Val Asn Ser
              85
Thr Glu Leu Asp Glu Leu Ile Gln Gln Ile Val Ala Ala Ile Asn Ala
100 105 110
                      105
Gly Ile Ile Pro Leu Gly Asn Thr Ser Asn Gln Ile Ser His Trp Asp
115 120 125
Leu Gly Ser Ser Phe Phe Phe Ala Gly Thr Val Ile Thr Thr Ile Gly
              135
 Phe Gly Asn Ile Ser Pro Arg Thr Glu Gly Gly Lys Ile Phe Cys Ile
                150
 Ile Tyr Ala Leu Leu Gly Ile Pro Leu Phe Gly Phe Leu Leu Ala Gly
                         170
 Val Gly Asp Gln Leu Gly Thr Ile Phe Gly Lys Gly Ile Ala Lys Val
                     185
 Glu Asp Thr Phe Ile Lys Trp Asn Val Ser Gln Thr Lys Ile Arg Ile
195 200 205
 Ile Ser Thr Ile Ile Phe Ile Leu Phe Gly Cys Val Leu Phe Val Ala
                      215
 Leu Pro Ala Ile Ile Phe Lys His Ile Glu Gly Trp Ser Ala Leu Asp
                          235
 Ala Ile Tyr Phe Val Val Ile Thr Leu Thr Thr Ile Gly Phe Gly Asp
245 250 255
                   230
  Tyr Val Ala Gly Gly Ser Asp Ile Glu Tyr Leu Asp Phe Tyr Lys Pro
260 265 270
  Val Val Trp Phe Trp Ile Leu Val Gly Leu Ala Tyr Phe Ala Ala Val
275 280 285
  Leu Ser Met Ile Gly Asp Trp Leu Arg Val Ile Ser Lys Lys Thr Lys
290 295 300
  Glu Glu Val Gly Glu Phe Arg Ala His Ala Ala Glu Trp Thr Ala Asn
320
  Val Thr Ala Glu Phe Lys Glu Thr Arg Arg Arg Leu Ser Val Glu Ile
                         330
  Tyr Asp Lys Phe Gln Arg Ala Thr Ser Ile Lys Arg Lys Leu Ser Ala
                               345
  Glu Leu Ala Gly Asn His Asn Gln Glu Leu Thr Pro Cys Arg Arg Thr
            340
                  360
  Leu Ser Val Asn His Leu Thr Ser Glu Arg Asp Val Leu Pro Pro Leu
370 375 380
   Leu Lys Thr Glu Ser Ile Tyr Leu Asn Gly Leu Thr Pro His Cys Ala
395 390 395 400
   Gly Glu Glu Ile Ala Val Ile Glu Asn Ile Lys
            405
         <210> 84
         <211> 20
         <212> DNA
```

<400> 84 catagccata ggtgaggact

<213> H. sapiens

<210> 85 <211> 20 <212> DNA

20

64

	PC	T/US99/03826
WO 99/43696		

<213> H. sapiens		
<400> 85		20
gagaggaaaa cagtctgggc		
<210> 86		•
<211> 20		
<212> DNA		
<213> H. sapiens		
<400> 86		20
ggacatcgaa ctaagacctg		
<210> 87		
<211> 20		
<212> DNA		
<213> H. sapiens		
<400> 87		20
tcccatgcca ttcagatctg		

International application No. PCT/US99/03826

		PC1/US99/03820
OF GUIDIECE VA	FTPD	
CLASSIFICATION OF SUBJECT MAPPC(6): C07H 21/04; C07K 14/705; C12N 1	5/09 15/63: C12O 1/68	
PC(6) :C07H 21/04; C07K 14/703; C12N 1 JS CL : 636/23.1, 24.3; 435/7.2, 69.1, 320.1	: 530/350	
JS CL: 636/23.1, 24.3; 435/7.2, 69.1, 320.1 cording to International Patent Classification (IPC) or to both national classification	n and IPC
THE DE CRADCHED		
inimum documentation searched (classification	system followed by classification sy	ymbols)
	; 530/350	
		the second of the fields searched
ocumentation searched other than minimum doc	amentation to the extent that such doc	Amend are mended in the Latest of
lectronic data base consulted during the intern	ational search (name of data base an	d, where practicable, search terms used)
	and the same of th	
Picase See Extra Sheet.		
	DEL PUANT	
DOCUMENTS CONSIDERED TO BE		Relevant to claim No.
	dication, where appropriate, of the rel	0720 7
The property of all	Cloning and Characteizati	on of a Novel 1-9
I T I Doctiti	mag Poraccillii Chaillei	110001111111111111111111111111111111111
Human Inward Recti	ine. FEBS Lett. 1998, Vol.	434, pages 171-
Expressed in Small Intest	ine. Peda Lea. 1996, 1986	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
176, see entire documen	. .	
		ł
		l l
		.
ļ		1
		Ì
	:	
·		
Further documents are listed in the co		patent family sonex.
Special entegories of cited documents:	dete sta	cument published after the international filing date or priority d not in conflict with the application but cited to understand ciple or theory underlying the invention
"A" document defining the general state of the art to be of particular relevance		
arre arrive document sublished on or after the is	sternational filing data conside	not of particular relevance; the considered to involve an inventive step- ared novel or estate the considered to involve an inventive step- the document is taken alone
of a document which may throw doubts on prior sited to establish the publication date of a	rity claim(s) or which is	to also also also also are als
Marie Marie (m sharen)	COEMO	and to severe all severes documents, such combination
O document referring to an oral disclosure,	being o	obvious to a person skilled in the wi
**Po document published prior to the international the priority date claimed	•	ent member of the same patent family
Date of the actual completion of the interna	tional search Date of mailing	g of the international search report 0 7 JUL 1999
28 MAY 1999		
Name of the ISA/US	Authorized off	Survence for
IVALUE BELL MARKET AND TO A Trademarks		1/ xumer (D)
Commissioner of Patents and Trademarks	NIRMAL	S. BASI
Box PCT Washington, D.C. 20231	NIRMAL Telephone No	

International application No. PCT/US99/03826

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, MEDLINE, JAPIO, BIOSIS, SCISEARCH, WPIDS, GENEMBL, NGENSEQ 34, EST, A-GENESEQ 32, PIR 58, SWISS-PROT 35, SPTREMBL 16. search terms: potassium channel, K+hnov

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO.2, the nucleic soid having the sequence of SEQ ID NO:1, nucleic soids hybridizing to said nucleic acids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of SEQ ID NO.2 and K+Hnov protein of SEQ ID NO.2.

Group II, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:4, the nucleic acid having the sequence of SEQ ID NO:3, nucleic acids hybridizing to said nucleic acids, express cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of SEQ ID NO:4 and K+Hnov protein of SEQ ID NO:4.

Group III, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:6, the nucleic acid having the sequence of SEQ ID NO:5, nucleic acids hybridizing to said nucleic acids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of SEQ ID NO:6 and K+Hnov protein of SEQ ID NO:6.

Group IV, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:8, the nucleic acid having the sequence of SEQ ID NO:7, nucleic acids hybridizing to said nucleic acids, expression cassette comprising said aucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of SEQ ID NO:8 and K+Haov protein of SEQ ID NO:8.

Group V, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:10, the nucleic acid having the sequence of SEQ ID NO:9, nucleic acids hybridizing to said nucleic acids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of

SEQ ID NO:10 and K+Hnov protein of SEQ ID NO:10. Group VI, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:12, the nucleic acid having the sequence of SEQ ID NO:11, nucleic acids hybridizing to said nucleic acids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing

K+Haov protein of SEQ ID NO:12 and K+Haov protein of SEQ ID NO:12. Group VII, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:14, the nucleic acid having the sequence of SEQ ID NO:13, nucleic acids hybridizing to said nucleic acids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Haov protein of SEQ ID NO:14 and K+Haov protein of SEQ ID NO:14.

Group VIII, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:16, the aucleic said having the sequence of SEQ ID NO:15, aucleic saids hybridizing to said nucleic saids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of SEQ ID NO:16 and K+Hnov protein of SEQ ID NO:16.

Group IX, claim(a)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:18, the aucleic acid having the sequence of SEQ ID NO:17, aucleic acids hybridizing to said nucleic acids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of SEQ ID NO:18 and K+Hnov protein of SEQ ID NO:18.

Group X, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:20, the aucleic acid having the sequence of SEQ ID NO:19, aucleic acids hybridizing to said nucleic acids, expression cassetts comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of SEQ ID NO:20 and K+Hnov protein of SEQ ID NO:20.

Group XI, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:25, the nucleic acid having the sequence of SEQ ID NO:21-25, nucleic acids hybridizing to said nucleic acids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of SEQ ID NO:25 and K+Hnov protein of SEQ ID NO:25.

Group XII, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:27, the nucleic acid having the sequence of SEQ ID NO:26, nucleic acids hybridizing to said nucleic acids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing

International application No. PCT/US99/03826

K+Hnov protein of SEQ ID NO:27 and K+Hnov protein of SEQ ID NO:27.

Group XIII, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:30, the nucleic acid having the sequence of SEQ ID NO:28-29, nucleic acids hybridizing to said nucleic acids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of SEQ ID NO:30 and K+Hnov protein of SEQ ID NO:30.

Group XIV, claim(s)1-9, draws to nucleic scids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:81, the nucleic acid having the sequence of SEQ ID NO:80, nucleic acids hybridizing to said nucleic acids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of SEQ ID NO:81 and K+Hnov protein of SEQ ID NO:81.

Group XV, claim(s)1-9, drawn to nucleic acids encoding K+Hnov protein having the amino acid sequence of SEQ ID NO:83, the nucleic acid having the sequence of SEQ ID NO:82, nucleic acids hybridizing to said nucleic acids, expression cassette comprising said nucleic acids, cell comprising said expression cassette, method for producing K+Hnov protein of SEQ ID NO:83 and K+Hnov protein of SEQ ID NO:83.

Group XVI, claim(s)10, draws to monoclonal antibody that binds to K+Hnov.

Group XVII, claim(s)11-14, drawn to non-human transgenic animal model for K+Hnov.

The inventions listed as Groups I-XVII do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: Group I is directed to nucleic acid (SEQ ID NO:1) encoding the K+Hnov protein of SEQ ID NO:2, nucleic acids hybridizing to said nucleic acid, expression cassette comprising said nucleic acid, cell comprising said cassette, method of producing the K+Hnov of SEQ ID NO:2 and the protein of SEQ ID NO:2. The special technical feature is the disclosed nucleic acid of SEQ ID NO:1 encoding the K+Hnov protein of SEQ ID NO:2. The aucleic acids, proteins, antibody and transgenic animal model of Groups II-XVII do not share the special technical feature of Group I wherein the products of said Groups are structurally and functionally different. As shown in Table 1, pages 8-9, the H+Nov proteins of SEQ ID NO: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 25, 27, 30, 81 and 83 are all structurally and functionally different, the nucleic acids encoding said proteins having different chromosome positions.

International application No. PCT/US99/03826

_		-		tions wher		rteia	claim	s wez	re fo	ound	luns	earc	chat	ble (C	Conti	nuat	ion	of item	1 of	first	sheet	1)			1
																									7
T	his in	tem	ational	report has t	not be	C2 C	tablish	od in	respo	ect o	of cer	tain (clain	ms un	der P	/Lacie	1/(.	ZXB) IOI	T O(U)	ULIOWI	ng ici	LIVIII D.			
1.]	Claim: bocau	s Nos.: se they rela	ute to	subj	oct ma	tter ne	ot re	niupe	red to	o be	scal	rched	by t	his A	utho	rity, n	mely	:					
2	. С]	hasan	s Nos.: so they rela tent that no	ste to j	perts ning:	of the	intern ernatio	natio onal	nal s naos	applic	catic an b	os th	aat do arriod	not:	comp speci	ly w ifical	ith the	presci	ibed :	requir	emen	ts to so	i ch	
3	· []	Claim	s Nos.: se they are	depen	rdent	claim	and a	are B	iot dr	rafted	l in a	acco	rdanc	e wit	h the	seco	nd and	third:	enten	ices of	f Rule	6.4(a)		
[Box I	u e	Observ	ations wh	ore u	mity	of in	rentio	मं क	laci	king	(Co	etir	nusti	D43 G	fiten	2 0	of first	shee)					4
ł				al Searchin																	VS:				ļ
				oe Extra Si																					ł
				JULIE 0																					
																				•					
	1.		As a	l required a	edditi	onal	search	foes v	were	o time	ely p	eid I	by ti	he ap	plica	nt, thi	is int	ematio	nal sc	arch I	eport	cove	ra all se	erchal	blo
	2.		As a	Il searchabl	le clai al fee	ims (could t	e scai	ırcho	d wi	ithou	t eff	iort j	ustify	ing s	n ad	ditio	sal foo,	this A	lutho	rity di	id not	invite	paym	cest
	3.		As o	aly some o those clair	f the i	roqui e wh	ired ad ich fee	dition:	re pe	parch aid, s	h food speci	s wei ifical	ro tin Lly c	moly daims	paid No	by th	ю вр	plicant,	this i	n terna	tions	l scar	ch rep	ort oov	023
	4.	x	rest	required ad icted to the EQ ID NO:	e invo	ntio	carch a first	focs V mentic	were	time dia	ely p	paid claim	by i	the a	pplic	ant. sd by	Con clai	sequen ms No	ily, ti	is in	ternati	ional	scarch	. герог	t is
	Ron	nari	k om F	rotest		7												e appl		prot	est.				